# HIV Molecular Immunology 2006/2007

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This publication is funded by the U.S. Department of Health and Human Services and the National Institutes of Health (Division of AIDS, National Institute of Allergy and Infectious Diseases) through an interagency agreement with the U.S. Department of Energy.



Published by
Theoretical Biology and Biophysics
Group T-10, Mail Stop K710
Los Alamos National Laboratory
Los Alamos, New Mexico 87545 U.S.A.

LA-UR 07-4752

http://www.hiv.lanl.gov/immunology



HIV Molecular Immunology 2006/2007

Published by Theoretical Biology and Biophysics Group T-10, Mail Stop K710 Los Alamos National Laboratory Los Alamos, New Mexico 87545 U.S.A

LA-UR 07-4752

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#### **Preface**

# Scope and purpose of the HIV molecular immunology database

HIV Molecular Immunology is a companion volume to HIV Sequence Compendium. This publication, the 2006/2007 edition, is the printed version of the web-based HIV Immunology Database (http://www.hiv.lanl.gov/content/immunology/). The web interface for this relational database has many search options, as well as interactive tools to help immunologists design reagents and interpret their results.

In the HIV Immunology Database, HIV-specific B-cell and T-cell responses are summarized and annotated. Immunological responses are divided into three parts, CTL, T helper, and antibody. Within these parts, defined epitopes are organized by protein and binding sites within each protein, moving from left to right through the coding regions spanning the HIV genome. We include human responses to natural HIV infections, as well as vaccine studies in a range of animal models and human trials. Responses that are not specifically defined, such as responses to whole proteins or monoclonal antibody responses to discontinuous epitopes, are summarized at the end of each protein section. Studies describing general HIV responses to the virus, but not to any specific protein, are included at the end of each part.

The annotation includes information such as cross-reactivity, escape mutations, antibody sequence, TCR usage, functional domains that overlap with an epitope, immune response associations with rates of progression and therapy, and how specific epitopes were experimentally defined. Basic information such as HLA specificities for T-cell epitopes, isotypes of monoclonal antibodies, and epitope sequences are included whenever possible. All studies that we can find that incorporate the use of a specific monoclonal antibody are included in the entry for that antibody. A single T-cell epitope can have multiple entries, generally one entry per study.

Finally, maps of all defined linear epitopes relative to the HXB2 reference proteins are provided. Alignments of CTL, helper T-cell, and antibody epitopes are available through the search interface on our web site at http://www.hiv.lanl.gov/content/immunology.

Only responses to HIV-1 and HIV-2 are included in the database. CTL responses to SIVs are periodically summarized in our review section by Dr. David Watkins and colleagues. (A review is included in this edition). Dr. Christian Brander and colleagues annually provide a concise listing of optimal CTL epitopes. Additional reviews that our editorial board deems of general interest to the HIV research immunology community are solicited each year. This year's reviews are printed in the first part of this database; reviews from previous years can be found at http://www.hiv.lanl.gov/content/hiv-db/REVIEWS/reviews.html.

Comments on the database or requests for the hard copy can be sent via email to immuno@lanl.gov.

#### Citing the database

This publication may be cited as

HIV Molecular Immunology 2006/2007. Bette T. M. Korber, Christian Brander, Barton F. Haynes, Richard Koup, John P. Moore, Bruce D. Walker, and David I. Watkins, editors. Publisher: Los Alamos National Laboratory, Theoretical Biology and Biophysics, Los Alamos, New Mexico.

#### About the cover

This year's cover illustrates the number of defined epitopes included in the database that span each position in the HIV proteome. The data are also plotted in Figure 1, which is shown on the next page. For T-cell epitopes, this density reflects the density of defined epitopes in the database, which in turn roughly reflects the density of responses detected against the whole genome by Elispots in natural infection. Both CD8+ T-cell (or cyototoxic T lymphocyte, maroon) and CD4+ T-cell (or Helper T cell, green) responses are most commonly detected in Gag and Nef, and we also have the highest density of database epitopes captured from the literature in these regions. In contrast, the antibody epitope database density in different locations (blue) is less meaningfully captured in this graph, because only continuous epitopes are included. Many antibody responses defined in the database are to discontinuous epitopes, or are defined regionally or by competition experiments, and other database entries are polyclonal responses with multiple antibodies binding to multiple regions; these not included in this map. The database entries have other biases in frequency. For example, the database is based on retrieval of information from the literature, and so a region like the V3 loop of HIV-1, which is of particular interest to many investigators, has been studied with great intensity, and this high level of interest accounts for the large spike of nearly 120 different antibody entries in gp120.

Preface About the PDF

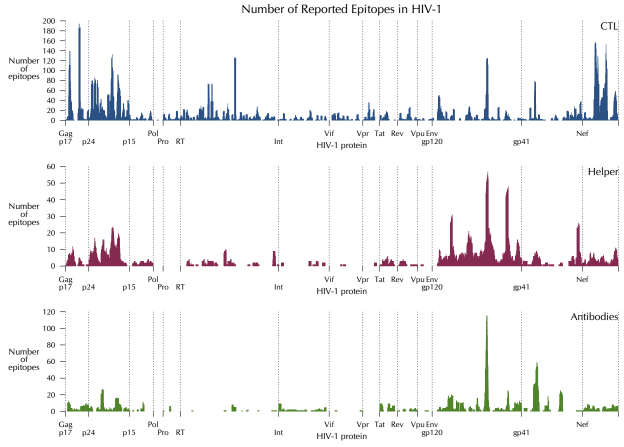
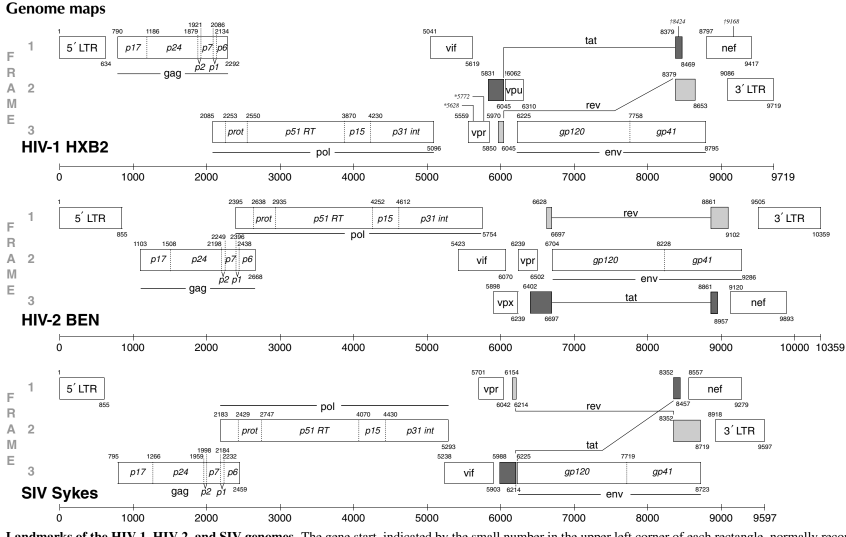


Figure 1: The number of defined epitopes included in the database that span each position in the HIV proteome.

#### **About the PDF**

The complete *HIV Molecular Immunology* 2006/2007 is available in Adobe Portable Document Format (PDF) from our website, http://www.hiv.lanl.gov/content/immunology. The PDF version is hypertext enabled and features 'clickable' table-of-contents, indexes, references and links to external web sites.

This volume is typeset using LATEX. The immunology data tables and epitope maps are produced automatically from the SQL database by a series of Perl programs.



Landmarks of the HIV-1, HIV-2, and SIV genomes. The gene start, indicated by the small number in the upper left corner of each rectangle, normally records the position of the a in the atg start codon for that gene while the number in the lower right records the last position of the stop codon. For *pol*, the start is taken to be the first t in the sequence ttttttag which forms part of the stem loop that potentiates ribosomal slippage on the RNA and a resulting -1 frameshift and the translation of the Gag-Pol polyprotein. The *tat* and *rev* spliced exons are shown as shaded rectangles. In HXB2, \*5628 and \*5772 mark positions of frameshifts in the *vpr* gene; !6062 indicates a defective acg start codon in *vpu*; †8424 and †9168 mark premature stop codons in *tat* and *nef*. See Korber *et al.*, Numbering Positions in HIV Relative to HXB2CG, in *Human Retroviruses and AIDS*, 1998, p. 102. Available from http://www.hiv.lanl.gov/content/hiv-db/REVIEWS/HXB2.html

Preface HIV/SIV proteins

#### **HIV/SIV** proteins

Name	Size	Function	Localization
Gag MA	p17	membrane anchoring; env interaction; nuclear transport of viral core (myristylated protein)	virion
CA	p24	core capsid	virion
NC	p7	nucleocapsid, binds RNA	virion
	p6	binds Vpr	virion
Protease (PR)	p15	gag/pol cleavage and maturation	virion
Reverse Transcriptase (RT)	p66, p51	reverse transcription	virion
RNase H	(heterodimer)	RNAse H activity	virion
Integrase (IN)		DNA provirus integration	virion
Env	gp120/gp41	external viral glycoproteins bind to CD4 and chemokine co-receptors	plasma membrane, virion envelope
Tat	p16/p14	viral transcriptional transactivator	primarily in nucleolus/nucleus
Rev	p19	RNA transport, stability and utilization factor (phosphoprotein)	primarily in nuleolus/nucleus shuttling between nucleolus and cytoplasm
Vif	p23	viral infectivity factor, inhibits minus-strand viral DNA hypermutation	cytoplasm (cytosol, membranes), virion
Vpr	p10-15	promotes nuclear localization of preintegration complex, inhibits cell division, arrests infected cells at G2/M	virion nucleus (nuclear membrane?)
Vpu	p16	promotes extracellular release of viral particles; degrades CD4 in the ER; (phosphoprotein only in HIV-1 and SIVcpz)	integral membrane protein
Nef	p27-p25	CD4 and class I downregulation (myristylated protein)	plasma membrane, cytoplasm, (virion?)
Vpx	p12-16	Vpr homolog present in HIV-2 and some SIVs, absent in HIV-1	virion (nucleus?)
Tev	p28	tripartite tat-env-rev protein (also named Tnv)	primarily in nucleolus/nucleus

**Abbreviations** Preface

#### **Abbreviations**

Common abbreviations and acronyms used in this database.

Abbrev.	Meaning				
AA	amino acid				
AAV	adeno-associated virus				
Ab	antibody				
ACTG	AIDS clinical trial group				
ADC	AIDS dementia complex				
ADCC	antibody-dependent cell-medicated				
	cytotoxicity				
ADE	antibody-dependent enhancement				
ADRA	Antiviral Drug Resistance Analysis: a				
	program that analyzes your sequences for				
	mutations known to confer drug resistance				
	and links to the records in the database				
AIDS	acquired immunodeficiency syndrome				
ANN	artificial neural networks				
anti MHC	anti major histocompatibility complex				
APC	antigen presenting cell				
ARC	AIDS related complex				
ART	anti-retroviral therapy				
AZT	azidothymidine				
BIMAS	•				
DIMAS	BioInformatics and Molecular Analysis				
DIV	Section				
BIV	bovine immunodeficiency virus				
BLAST	Basic Local Alignment Search Tool				
CAEV	caprine arthritis/encephalitis virus				
CD4BS	CD4 binding site				
CD4i	antibody that has enhanced binding to gp120				
~~ ~	in the presence of SCD4 (CD4 induced)				
CDC	Centers for Disease Control and Prevention				
CDR	complementary determining regions				
CFA	complete Freund's adjuvant				
CHI	Center for HIV Information				
CMI	cell-mediated immunity				
CMV	cytomegaliovirus				
CNS	central nervous system				
CP	canary pox				
CRF	circulating recombinant form				
CsA	cyclosporine A				
CSF	cerebrospinal fluid				
CTL	cytotoxic T lymphocyte				
CTLe	CTL effector				
CTLp	CTL precursor				
CyPA	cyclophilin A				
DC	dendritic cell				
DDDP	DNA-dependent DNA polymerase				
DHH	U. S. Department of Health and Human				
	Services				
dMM	deopymannojirimycin				
dpc	days post challenge				
DTT	dithiothrietol				
עוע	ummounictor				

Abbrev.	Meaning	
EIA	enzyme immuno assay	
EIAV	equine infectious anemia virus	
ELF	Epitope Location Finder	
ELISA	Enzyme Linked ImmunoSorbent Assay	
ER	endoplasmic reticulum	
Fabs	fragment antigen binding-univalent	
	antibody fragment	
FATT-CTL	Fluorescent antigen-transfected target	
	cell–CTL	
FIV	feline immunodeficiency virus	
FP	fowl pox	
FSW	female sex worker	
GALT	gut-associated lymphoid tissues	
GDE format	Genetic Data Environment	
gp	glycoprotein	
GRIV	genetic resistance to HIV	
HAART	highly-active anti-retroviral therapy	
HCV	hepatitis C virus	
HEPS	HIV-exposed persistently seronegative	
HIV	human immunodeficiency virus	
HIVD	HIV-1 dementia	
HLA	human leukocyte antigens	
HLA-MHC	human leukocyte antigens-major	
IILA-WIIIC	histocompatibility complex	
HMM	hidden Markov models	
IAVI	International AIDS Vaccine Initiative	
IDE		
	immunodominant epitope	
IE genes IFA	immediate early genes	
IFA IFN	incomplete Freund's adjuvant interferon	
IGN IG format	IntelliGenetics format	
Ig 11	immunoglobulin interleukin	
IL mu	11100110011111	
INHI :	immunologically normal HIV-infected	
iscom	immunostimulating complex	
KLH	keyhole limpet hemocyanin	
LANL	Los Alamos National Laboratory	
LDA	limiting dilution assay	
LN	lymph node	
LPR	lymphoproliferative response	
LT	labile enterotoxin	
LTNP	long-term non-progressor	
LTR	long terminal repeat	
LTS	long term survivor	
mAb	monoclonal antibody	
MBL	mannose-binding lectin	
MCMC	Markov chain Monte Carlo	
MDP	muramyl dipeptide	
MEI	multiple epitope immunogen	
MHC	major histocompatibility complex	
MHR	major homology region	
ML	maximum likelihood	
MLV	murine leukemia virus	

Preface Amino Acid Codes

Abbrev.	Meaning			
MP	maximum parsimony			
mpc	months post challenge			
MPER	membrane-proximal external region			
MRC	Medical Research Council, UK			
MSF	multiple sequence alignment format of the			
	GCG sequence analysis package			
MV	measles vector			
MVA vector	modified vaccinia virus Ankara			
Nab	neutralizing antibody			
NCBI	National Center for Biotechnology			
	Information			
NIAID	National Institute of Allergies and			
	Infectious Diseases			
NIBSC	National Institute for Biological Standards			
	and Control, UK			
NIH	National Institutes of Health			
NIST	National Institute of Standards and			
	Technology			
NJ	neighbor joining			
NLS	nuclear localization signal			
NRP	non-rapid progressor			
NSI	non-synctium-inducing			
p	protein			
PB	peripheral blood			
PBL	peripheral blood lymphocyte			
PBMC	peripheral blood mononuclear cell			
PCOORD	principal coordinate analysis			
PCR	polymerase chain reaction			
PERV	porcine endogenous retrovirus			
PHYLIP	Phylogeny Inference Package			
PL	proteoliposome			
RAC	ricin A chain			
RDDP	RNA-dependent DNA polymerase			
rec/r	recombinant			
RIP	Recombinant Identification Program: a			
	program for detecting evidence of			
	inter-subtype recombination			
RIPA	Radio Immuno Precipitation Assay			
RP	rapid progressor			
RRE	Rev-responsive element			
rsgp160	recombinant soluble gp160			
RSV	Rous sarcoma virus			
SAM	Sequence Alignment and Modeling			
	program			
SAP	sequential antigen panning			
sCD4	soluble CD4			
scFv	single-chain variable fragment			
SDS	sodium duodecyl sulfate			
SFV	Semliki Forest virus			
SI	synctium inducing			
SIV	simian immunodeficiency virus			

Abbrev.	Meaning
SIVE	SIV encephalitis
SLE	systemic lupus erythhermatosis
SNAP	synonymous-nonsynonymous analysis
	program
STI	supervised treatment interruption (also seen as
	structured treatment interruption and standard
	treatment interruption)
TCLA	T cell line adapted
TCR	T-cell receptor
Th	T-helper cell
TNF	tumor necrosis factor
VEE	Venezuelan equine encephalitis
<b>VESPA</b>	Viral Epidemiology Signature Pattern Analysis
VIP	vasoactive intestinal peptide
VL	viral load
VLP	virus like particle, assembled from p55 gag
VSV	vesicular stomatitis virus
VV	vaccinia virus
WB	Western Blot

#### **Amino Acid Codes**

Α	Alanine
В	Aspartic Acid or Asparagine
C	Cysteine
D	Aspartic Acid
Ε	Glutamic Acid
F	Phenylalanine
G	Glycine
Н	Histidine
Ι	Isoleucine
K	Lysine
L	Leucine
М	Methionine
Ν	Asparagine
Р	Proline
Q	Glutamine
R	Arginine
S	Serine
Т	Threonine
٧	Valine
W	Tryptophan
Χ	unknown or "other" amino acid
Υ	Tyrosine
Z	Glutamic Acid or Glutamine
	gap
-	identity
\$	stop codon
#	frameshift

# Part I Review Articles

# Identification of HIV-Derived, HLA Class I Restricted CTL Epitopes: Insights into TCR Repertoire, CTL Escape and Viral Fitness

Nicole Frahm<sup>a</sup>, Caitlyn Linde<sup>a</sup>, Christian Brander<sup>a</sup>

# I-A-1 The importance of well-defined T cell epitopes in understanding host immunity to HIV

A detailed understanding of T cell immunity to HIV infection will be required for the design and development of an effective HIV vaccine. Over the last few years, it has become clear that the mere breadth and magnitude of T cell responses directed against the entire viral proteome are not associated with immune control and that a more in-depth look at T cell specificity, effector functions and viral diversity will be needed to define true correlates of immune protection [Zuñiga et al., 2006; Frahm et al., 2004; Kiepiela et al., 2004; Betts et al., 2006; Masemola et al., 2004a]. In particular, the relationship between targeting specific regions of the viral genome, T cell escape and, as a consequence, changes in viral replicative fitness has become a focus of much debate [Zuñiga et al., 2006; Masemola et al., 2004a; Martinez-Picado et al., 2006; Bailey et al., 2006; Li et al., 2007; Liu et al., 2006; Yeh et al., 2006; Ganusov & De Boer, 2006]. In addition, studies on both the transmission and reversion of CTL escape variants, the induction of T cell specificities against effective viral escape variants as well as work addressing the role of subdominant T cell responses in the control of HIV have provided a better understanding of the complex dynamics between host immune response and viral adaptation to immune pressure [Leslie et al., 2004, 2005; Friedrich et al., 2004; Allen et al., 2005b,a; Frahm et al., 2006a]. For most of these studies, the identification of precisely defined HLA class I-restricted CTL epitopes has been key and will continue to be a central prerequisite, especially in

studies that focus on less well studied human populations with diverse HLA backgrounds [Kiepiela *et al.*, 2004].

# I-A-2 Escape pathways of dominant CTL epitopes

Despite the increasing appreciation for the role of subdominant CTL responses in viral control, much of the current knowledge on immune driven viral evolution and CTL escape is based on the study of a few, well defined, dominant T cell targets [Bailey et al., 2006; Leslie et al., 2004, 2005; Frahm et al., 2006a; Brander et al., 1998; Iversen et al., 2006; Migueles et al., 2003; Yu et al., 2006]. In some of these cases, for instance the dominant HLA-B27-restricted epitope KK10 in HIV Gag p24 (KRWIILGLNK), CTL escape from a single epitope can result in elevated viral loads and accelerated disease progression [Goulder et al., 1997c; Feeney et al., 2004]. However, for other epitopes and alleles, the relationship between CTL escape and disease progression may be more complex. For instance, in the case of the HLA-B57-restricted KF11 epitope (KAFSPEV-IPMF), a number of studies have found intact viral epitope sequences and significant epitope-specific responses even in individuals with seemingly uncontrolled viral replication [Migueles et al., 2003]. On the other hand, others have reported "escape" mutations in the KF11 epitope in individuals with elite control of viral replication [Bailey et al., 2006], indicating that additional factors are likely crucially involved in shaping an effective T cell response to this virus. Indeed, recent data generated by multi-color flow cytometric analyses highlight the importance of polyfunctional effector cells in the control of HIV. These polyfunctional T cells may be impaired in some individuals with progressive disease, although these subjects may have conserved epitope sequences and strong, epitope-specific T cell responses as assessed by interferon-γ secretion [Betts et al., 2006].

A number of reports have now also begun to address the kinetics of compensatory mutations that are either required for effective T cell escape or for the maintenance of viral replicative fitness [Yeh *et al.*, 2006; Iversen *et al.*, 2006; Kelleher *et al.*, 2001; Peyerl *et al.*, 2004]. Recent

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In *HIV Molecular Immunology 2006/2007*. Bette T. M. Korber, Christian Brander, Barton F. Haynes, Richard Koup, John P. Moore, Bruce D. Walker, and David I. Watkins, editors. Publisher: Los Alamos National Laboratory, Theoretical Biology and Biophysics, Los Alamos, New Mexico. p. 3–28.

analyses by Iversen et al. [2006] have addressed viral escape pathways in the context of the dominant HLA-A2 restricted epitope SLYNTVATL in HIV Gag p17, showing that effective viral escape was only achieved after serial mutations in three positions within the optimal epitope. These changes were required for escape from TCR recognition, indicating that simple reduction of epitope binding affinity to the presenting HLA class I molecule does not necessarily allow the virus to evade immune control. Rather, effective escape may occur only when there is a profoundly diminished interaction between TCR and the HLA/peptide complex, to which the epitope binding affinity will only partially contribute. This highlights the need to expand the analyses of HIV-specific immune responses to include detailed assessments of functional avidity of these responses as well as to consider the impact that the T cell receptor repertoire diversity may have on the emergence of CTL escape variants. Thus, a number of studies have started to shed some light on the factors that define a broadly cross-reactive and highly avid T cell response, which may be especially well suited to prevent viral escape. Of note, a number of recent analyses have indicated that a more narrow T cell receptor repertoire, mediating T cell responses of high functional avidity, may be particularly effective in controlling viral replication [Yu et al., 2006; Price et al., 2004; Frahm et al., 2006b; Ahlers et al., 2001; Alexander-Miller et al., 1996a]. Although one might expect a broader epitope-specific TCR repertoire to provide more possible candidate TCR to effectively recognize emerging escape variants, our own data and studies from other laboratories indicate that an increased breath of the TCR repertoire may be associated with reduced functional avidity of the total epitope-specific T cell population [Yu et al., 2006; Price et al., 2004; Frahm et al., 2006b; Messaoudi et al., 2002]. This is highlighted in a recent analysis of the TCR usage in T cells directed against the dominant HIV Gag KF11 (KAFSPEVIPMF) epitope, which can be restricted by the two closely related alleles HLA-B\*5701 and B\*5703. In the context of HLA-B\*5701, the KF11 specific response is characterized by a TCR repertoire that is highly conserved among HLA-B\*5701-expressing individuals and that efficiently cross-reacts with viral epitope variants [Yu et al., 2006]. In contrast, in the context of HLA-B\*5703, the KF11 epitope induces an entirely different, more heterogeneous TCR repertoire that fails to recognize the most frequently occurring epitope variants, indicating that extensive TCR diversity may not effectively prevent the emergence of epitope escape variants. How the immune system may control the emergence of highly avid rather than broad TCR repertoires may at least partly depend on antigen availability and competition of expanding T cell populations [Kim et al., 2006; Price et al., 2005]. Studies that address these factors in HIV and other viral infections will have obviously important implications for vaccine design, which may need to consider different lev-

els of antigen availability during the induction phase of responses and aim to specifically drive the expansion of high avidity T cell responses of limited TCR diversity and (thus) superior ability to recognize epitope variants.

# I-A-3 Inclusion of epitopes in the optimal list

As every year, the present listing is based on the inclusion of epitopes that fulfill a number of stringent criteria intended to ensure reliable description of the optimal length epitope and correct assignment of the restricting HLA class I allele(s) [Hunziker et al., 1998; Brander & Walker, 1995]. Nevertheless, there may still be occasions where the data reported here conflict with data in other laboratories and we to encourage any investigators who observe discrepancies in their own data and what is reported here to bring this to our attention. The selection of inclusion criteria itself is obviously subject to potential disagreement, too. In particular, a number of epitopes have been described over the last year where binding motif algorithms have been used to predict the optimal epitope length, or for which the restricting HLA class I alleles have been inferred by statistical associations or binding assays [Kawashima et al., 2005; Satoh et al., 2005]. Although in some large cohort analyses, a number of associations have reached statistically highly significant associations, the optimal epitope was often inferred based on previously published motif data [Kiepiela et al., 2004; Honeyborne et al., 2006]. We have thus opted not include these epitopes even though a number of them are likely to be accurately described. However, as binding motif predictions are only as good as the quality of the described epitopes for a given allele, care must be taken to not further "confirm" existing binding motifs by data that were generated based on the original training set defining the motifs in the first place. Thus, at least some of the inferred optimal epitope lengths should ideally be confirmed experimentally to ensure that sometimes only loosely defined allele-specific binding motifs are indeed correct. Thus, while newly identified epitopes may violate known HLA binding motifs, these extensively mapped epitopes may help to refine currently incompletely defined allele-specific binding motifs. The expansion of binding motifs to include less frequently used amino acids will not only facilitate work in the HIV field, but also in other viral infections, cancer and autoimmunity.

#### I-A-4 Table of optimal HIV-1 CTL epitopes

 Table I-A.1: Best defined HIV CTL epitopes.

HLA	Protein	AA	Sequence	Reference
A*0101 (A1)	gp160	787–795	RRGWEVLKY	Cao, 2002
A2	RT	127–135	YTAFTIPSV	Draenert, 2004
A*0201 (A2)		1° ancl	2 6 C nor <b>L L</b>	Falk et al., 1991; Barouch et al., 1995
		2° ancl	nor V	
A*0201 (A2)	p17	77–85	SLYNTVATL	Johnson et al., 1991; Parker et al., 1992, 1994
A*0201 (A2)	p2p7p1p6	70–79	FLGKIWPSYK	Yu et al., 2002b
A*0201 (A2)	Protease	76–84	LVGPTPVNI	Karlsson et al., 2003
A*0201 (A2)	RT	33-41	ALVEICTEM	Haas et al., 1998; Haas, 1999
A*0201 (A2)	RT	179–187	VIYQYMDDL	Harrer et al., 1996a
A*0201 (A2)	RT	309-317	ILKEPVHGV	Walker et al., 1989; Tsomides et al., 1991
A*0201 (A2)	Vpr	59–67	AIIRILQQL	Altfeld et al., 2001a,b
A*0201 (A2)	gp160	311-320	RGPGRAFVTI	Alexander-Miller et al., 1996b
A*0201 (A2)	gp160	813-822	SLLNATDIAV	Dupuis <i>et al.</i> , 1995
A*0201 (A2)	Nef	136-145	PLTFGWCYKL	Haas et al., 1996; Maier & Autran, 1999
A*0201 (A2)	Nef	180–189	VLEWRFDSRL	Haas et al., 1996; Maier & Autran, 1999
A*0202 (A2)			2 C L L V	Barouch et al., 1995
A*0202 (A2)	p17	77–85	SLYNTVATL	Goulder, 1999
A*0205 (A2)	p17	77–85	SLYNTVATL	Goulder, 1999
A*0205 (A2)	gp160	846-854	RIRQGLERA	Sabbaj <i>et al.</i> , 2003
A*0205 (A2)	Nef	83–91	GAFDLSFFL	Rathod, 2006
A*0207 (A2)	p24	164–172	YVDRFYKTL	Currier et al., 2002

 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
A*0301 (A3)			2 C	Marsh et al., 2000
			L K	
			V Y	
			М	
			I F	
			A R	
			<u>S</u>	
		10.55	T	
A*0301 (A3)	p17	18–26	KIRLRPGGK	Harrer <i>et al.</i> , 1996b
A*0301 (A3)	p17	20–28	RLRPGGKKK	Goulder <i>et al.</i> , 1997b; Culmann, 1999; Lewinsohn & Riddell, 1999; Wilkes & Ruhl, 1999
A*0301 (A3)	p17	20–29	RLRPGGKKKY	Goulder et al., 2000b
A*0301 (A3)	RT	33–43	ALVEICTEMEK	Haas et al., 1998; Haas, 1999
A*0301 (A3)	RT	73–82	KLVDFRELNK	Yu et al., 2002a
A*0301 (A3)	RT	93–101	GIPHPAGLK	Yu <i>et al.</i> , 2002a
A*0301 (A3)	RT	158–166	AIFQSSMTK	Threlkeld et al., 1997
A*0301 (A3)	RT	269–277	QIYPGIKVR	Yu et al., 2002a
A*0301 (A3)	RT	356–366	RMRGAHTNDVK	Yu et al., 2002a
A*0301 (A3)	Integrase	179–188	AVFIHNFKRK	Yu et al., 2002a
A*0301 (A3)	Vif	17–26	RIRTWKSLVK	Altfeld et al., 2001a; Yu et al., 2002a
A*0301 (A3)	Vif	28–36	HMYISKKAK	Yu et al., 2002a
A*0301 (A3)	Vif	158–168	KTKPPLPSVKK	Yu et al., 2002a
A*0301 (A3)	Rev	57–66	ERILSTYLGR	Addo, 2002; Yu <i>et al.</i> , 2002a
A*0301 (A3)	gp160	37–46	TVYYGVPVWK	Johnson <i>et al.</i> , 1994
A*0301 (A3)	gp160	770–780	RLRDLLLIVTR	Takahashi <i>et al.</i> , 1991
A*0301 (A3) A*0301 (A3)	Nef Nef	73–82 84–92	QVPLRPMTYK AVDLSHFLK	Koenig <i>et al.</i> , 1990; Culmann <i>et al.</i> , 1991 Yu <i>et al.</i> , 2002a
	1101	01 72		
A*1101 (A11)			2 C <b>K</b>	Zhang et al., 1993; Rammensee et al., 1995
			V	
			Ī	
			- F	
			Y	
A*1101 (A11)	p17	84–92	TLYCVHQRI	Harrer et al., 1998
A*1101 (A11)	p24	217-227	ACQGVGGPGHK	Sipsas <i>et al.</i> , 1997
A*1101 (A11)	RT	158-166	AIFQSSMTK	Johnson & Walker, 1994; Zhang <i>et al.</i> , 1993;
				Threlkeld et al., 1997
A*1101 (A11)	RT	341-350	IYQEPFKNLK	Culmann, 1999
A*1101 (A11)	RT	520-528	QIIEQLIKK	Fukada <i>et al.</i> , 1999
A*1101 (A11)	Integrase	179–188	AVFIHNFKRK	Fukada et al., 1999
A*1101 (A11)	gp160	199–207	SVITQACPK	Fukada et al., 1999
A*1101 (A11)	Nef	73–82	QVPLRPMTYK	Buseyne, 1999
A*1101 (A11)	Nef	75–82	PLRPMTYK	Culmann et al., 1991
A*1101 (A11)	Nef	84–92	AVDLSHFLK	Culmann et al., 1991
A23	gp160	585-593	RYLKDQQLL	Cao et al., 2003

 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
A*2402 (A24)			2 C	Maier et al., 1994
			ΥI	
			L F	
A*2402 (A24)	p17	28–36	KYKLKHIVW	Ikeda-Moore et al., 1998; Lewinsohn, 1999
A*2402 (A24)	p24	162–172	RDYVDRFFKTL	Dorrell et al., 1999; Rowland-Jones, 1999
A*2402 (A24)	gp160	52-61	LFCASDAKAY	Lieberman et al., 1992; Shankar et al., 1996
A*2402 (A24)	gp160	585-593	RYLKDQQLL	Dai <i>et al.</i> , 1992
A*2402 (A24)	Nef	134–141	RYPLTFGW	Goulder et al., 1997a; Ikeda-Moore et al., 1998
A*2501 (A25)	p24	13–23	QAISPRTLNAW	Kurane & West, 1999
A*2501 (A25)	p24	71–80	ETINEEAAEW	Klenerman et al., 1996; van Baalen et al., 1996
A*2501 (A25)	gp160	703–712	EIIFDIRQAY	Liu et al., 2006
A*2601 (A26)			12 6 C	Dumrese et al., 1998
` ,			V Y	,
			T F	
			I	
			L	
			F	
			D I	
			E L V	
A*2601 (A26)	p24	35–43	EVIPMFSAL	Goulder et al., 1996a
A*2601 (A26)	RT	449–457	ETKLGKAGY	Sabbaj et al., 2003
A29	Nef	120–128	YFPDWQNYT	Draenert et al., 2004
A*2902 (A29)	p17	78–86	LYNTVATLY	Masemola et al., 2004b
A*2902 (A29)	gp160	209–217	SFEPIPIHY	Altfeld, 2000
. ,				
A30	p17	34–44	LVWASRELERF	Masemola et al., 2004b
A*3002 (A30)			12 C	Rammensee et al., 1999
			Y Y	
			F	
			V	
			<b>v</b> R	
A*3002 (A30)	p17	76–86	RSLYNTVATLY	Goulder et al., 2001
A*3002 (A30)	RT	173–181	KQNPDIVIY	Goulder et al., 2001 Goulder et al., 2001
A*3002 (A30)	RT	263–271	KLNWASQIY	Goulder et al., 2001
A*3002 (A30)	RT	356–365	RMRGAHTNDV	Sabbaj <i>et al.</i> , 2003
A*3002 (A30)	Integrase	219–227	KIQNFRVYY	Sabbaj <i>et al.</i> , 2003; Rodriguez <i>et al.</i> , 2004
A*3002 (A30)	gp160	310-318	HIGPGRAFY	Sabbaj <i>et al.</i> , 2003
A*3002 (A30)	gp160	704–712	IVNRNRQGY	Goulder et al., 2001
A*3002 (A30)	gp160	794-802	KYCWNLLQY	Goulder et al., 2001

 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
A*3101 (A31)			2 C	Falk et al., 1994; Rammensee et al., 1999
			R	
			L	
			V	
			Υ	
			F	
A*3101 (A31)	gp160	770–780	RLRDLLLIVTR	Safrit et al., 1994a,b
A*3201 (A32)	RT	392-401	PIQKETWETW	Harrer et al., 1996b
A*3201 (A32)	gp160	419–427	RIKQIINMW	Harrer et al., 1996b
A33	Nef	133–141	TRYPLTFGW	Cao, 2002
A*3303 (A33)	gp160	698–707	VFAVLSIVNR	Hossain et al., 2001
A*3303 (A33)	gp160	831-838	EVAQRAYR	Hossain et al., 2001
A*3303 (A33)	Vpu	29–37	EYRKILRQR	Addo et al., 2002
A66	RT	438–448	ETFYVDGAANR	Rathod, 2006
A*6801 (A68)	Tat	39–49	ITKGLGISYGR	Oxenius et al., 2002
A*6801 (A68)	Vpr	52–62	DTWAGVEAIIR	Sabbaj <i>et al.</i> , 2004
A*6802 (A68)	RT	436–445	GAETFYVDGA	Rathod & Kiepiela, 2005
A*6802 (A68)	Protease	3–11	ITLWQRPLV	Rowland-Jones, 1999
A*6802 (A68)	Protease	30-38	DTVLEEWNL	Rowland-Jones, 1999
A*6802 (A68)	Vpr	48-57	ETYGDTWTGV	Rathod & Kiepiela, 2005
A*6802 (A68)	gp160	777–785	IVTRIVELL	Wilkes, 1999
A*7401 (A19)	Protease	3–11	ITLWQRPLV	Rowland-Jones, 1999

 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
B7	p24	84–92	HPVHAGPIA	Yu <i>et al.</i> , 2002a
B7	RT	156-164	SPAIFQSSM	Linde & Faircloth, 2006
B7	Rev	41–50	RPAEPVPLQL	Yang, 2006
B*0702 (B7)			123 C	Englehard et al., 1993; Rammensee et al., 1999
			P L	
			AR <b>F</b>	
			RK	
B*0702 (B7)	p24	16–24	SPRTLNAWV	Lewinsohn, 1999
B*0702 (B7)	p24	48–56	TPQDLNTML	Wilson, 1999; Wilkes <i>et al.</i> , 1999; Jin <i>et al.</i> , 2000; Wilson <i>et al.</i> , 1997
B*0702 (B7)	p24	223-231	GPGHKARVL	Goulder, 1999
B*0702 (B7)	Vpr	34-42	FPRIWLHGL	Altfeld et al., 2001a
B*0702 (B7)	Vif	48-57	HPRVSSEVHI	Altfeld et al., 2001a
B*0702 (B7)	gp160	298-307	RPNNNTRKSI	Safrit <i>et al.</i> , 1994b
B*0702 (B7)	gp160	843-851	IPRRIRQGL	Wilkes & Ruhl, 1999
B*0702 (B7)	Nef	68–77	FPVTPQVPLR	Haas et al., 1996; Maier & Autran, 1999
B*0702 (B7)	Nef	68–76	FPVTPQVPL	Bauer et al., 1997; Frahm & Goulder, 2002
B*0702 (B7)	Nef	71–79	TPQVPLRPM	Goulder, 1999
B*0702 (B7)	Nef	77–85	RPMTYKAAL	Bauer et al., 1997
B*0702 (B7)	Nef	106-115	RQDILDLWIY	Goulder, 1999
B*0702 (B7)	Nef	128-137	TPGPGVRYPL	Culmann-Penciolelli et al., 1994; Haas et al., 1996
B8	gp160	848–856	RQGLERALL	Cao, 2002
B*0801 (B8)			23 5 C	Hill et al., 1992; Sutton et al., 1993; DiBrino et al., 1994b
			K K L	,
			R	
			PR L	
B*0801 (B8)	p17	24–32	GGKKKYKLK	Reid et al., 1996; Goulder et al., 1997d
B*0801 (B8)	p17	74–82	ELRSLYNTV	Goulder <i>et al.</i> , 1997d
B*0801 (B8)	p24	128–135	EIYKRWII	Sutton <i>et al.</i> , 1993; Goulder <i>et al.</i> , 1997d
B*0801 (B8)	p24	197–205	DCKTILKAL	Sutton <i>et al.</i> , 1993
B*0801 (B8)	RT	18-26	GPKVKQWPL	Walker et al., 1989; Sutton et al., 1993
B*0801 (B8)	gp160	2-10	RVKEKYQHL	Sipsas <i>et al.</i> , 1997
B*0801 (B8)	gp160	586-593	YLKDQQLL	Johnson <i>et al.</i> , 1992; Shankar <i>et al.</i> , 1996
B*0801 (B8)	Nef	13-20	WPTVRERM	Goulder et al., 1997d
B*0801 (B8)	Nef	90–97	FLKEKGGL	Culmann-Penciolelli et al., 1994; Price et al.,
,				1997
B13	p24	3–11	VQNLQGQMV	Honeyborne et al., 2007
B13	p24	94–104	GQMREPRGSDI	Honeyborne et al., 2007
B13	p2p7p1p6	66–74	RQANFLGKI	Honeyborne et al., 2007
B13	Protease	57–66	RQYDQILIEI	Honeyborne et al., 2007
B13	RT	333–341	GQGQWTYQI	Honeyborne et al., 2007
B13	Nef	106–114	RQDILDLWI	Harrer et al., 2005
B13	Nef	106–114	RQDILDLWV	Honeyborne et al., 2007

 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
B14	p2p7p1p6	42–50	CRAPRKKGC	Yu et al., 2002b
B*1401 (B14)	RT	142–149	IRYQYNVL	Rathod, 2006
B*1402 (B14)			23 5 C <b>R R L K H</b> L  Y  F	DiBrino et al., 1994a
B*1402 (B14)	p24	166-174	DRFYKTLRA	Harrer et al., 1996b
B*1402 (B14)	gp160	584–592	ERYLKDQQL	Johnson et al., 1992
B*1501 (B62)			2 C	
			Q Y	Barber <i>et al.</i> , 1997
			L F	Barber <i>et al.</i> , 1997
			M	Barber <i>et al.</i> , 1997
B*1501 (B62)	p24	137–145	GLNKIVRMY	Johnson et al., 1991; Goulder, 1999
B*1501 (B62)	RT	260–271	LVGKLNWASQIY	Johnson, 1999
B*1501 (B62)	RT	309–318	ILKEPVHGVY	Johnson et al., 1991; Johnson, 1999
B*1501 (B62)	Nef	117–127	TQGYFPDWQNY	Culmann, 1999
B*1503 (B72)	p24	24-32	VKVIEEKAF	Honeyborne & Kiepiela, 2005
B*1503 (B72)	p24	164–172	YVDRFFKTL	Masemola et al., 2004b
B*1503 (B72)	Protease	68–76	GKKAIGTVL	Rathod & Bishop, 2006
B*1503 (B72)	RT	496–505	VTDSQYALGI	Sabbaj et al., 2003
B*1503 (B72)	Integrase	135–143	IQQEFGIPY	Honeyborne & Kiepiela, 2005
B*1503 (B72)	Integrase	185–194	FKRKGGIGGY	Honeyborne, 2003
B*1503 (B72)	Integrase	263–271	RKAKIIRDY	Cao et al., 2003
B*1503 (B72)	Tat	38–47	FQTKGLGISY	Novitsky et al., 2001
B*1503 (B72)	Nef	183–191	WRFDSRLAF	Cao, 2002
B*1510 (B71)	p24	12-20	HQAISPRTL	Day, 2005
B*1510 (B71)	p24	61–69	GHQAAMQML	Day, 2003
B*1510 (B71)	Integrase	66–74	THLEGKIIL	Kiepiela et al., 2007
B*1510 (B71)	Vif	79–87	WHLGHVSI	Honeyborne, 2003
B*1516 (B63)			2 9 T Y S I V	Barber et al., 1997; Seeger et al., 1998
B*1516 (B63)	gp160	375–383	<b>F</b> SFNCGGEFF	Wilson et al., 1997; Wilson, 1999

 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
B18	RT	137–146	NETPGIRYQY	Rathod & Bishop, 2006
B18	RT	175–183	NPEIVIYQY	Rathod, 2006
B18	Nef	105–115	RRQDILDLWVY	Yang, 2006
B*1801 (B18)	p24	161–170	FRDYVDRFYK	Ogg et al., 1998
B*1801 (B18)	Vif	102-111	LADQLIHLHY	Altfeld et al., 2001a
B*1801 (B18)	gp160	31–39	AENLWVTVY	Liu et al., 2006
B*1801 (B18)	gp160	61–69	YETEVHNVW	Liu <i>et al.</i> , 2006
B*1801 (B18)	Nef	135–143	YPLTFGWCY	Culmann <i>et al.</i> , 1991; Culmann-Penciolelli <i>et al.</i> , 1994
B27	Vpr	31–39	VRHFPRIWL	Addo & Rathod, 2004
B*2703 (B27)	p24	131–140	RRWIQLGLQK	Rowland-Jones et al., 1998; Rowland-Jones, 1999
B*2705 (B27)			12 C R L F	Jardetzky et al., 1991; Rammensee et al., 1995
			К К	
			R R	
			G I	
			A	
B*2705 (B27)	p17	19–27	IRLRPGGKK	McKinney et al., 1999; Lewinsohn, 1999
B*2705 (B27)	p24	131–140	KRWIILGLNK	Nixon <i>et al.</i> , 1988; Buseyne <i>et al.</i> , 1993; Goulder <i>et al.</i> , 1997c
B*2705 (B27)	gp160	786–795	GRRGWEALKY	Lieberman et al., 1992; Lieberman, 1999
B*2705 (B27)	Nef	105–114	RRQDILDLWI	Goulder et al., 1997b
B*3501 (B35)			2 C	Hill et al., 1992; Rammensee et al., 1999
			P Y	
			A F	
			V M	
			S L I	
B*3501 (B35)	p17	36-44	WASRELERF	Goulder et al., 1997a
B*3501 (B35)	p17	124-132	NSSKVSQNY	Rowland-Jones et al., 1995
B*3501 (B35)	p24	122-130	PPIPVGDIY	Rowland-Jones et al., 1995
B*3501 (B35)	p24	122-130	NPVPVGNIY	Rowland-Jones et al., 1995
B*3501 (B35)	RT	107-115	TVLDVGDAY	Wilkes & Ruhl, 1999; Wilson et al., 1999
B*3501 (B35)	RT	118-127	VPLDEDFRKY	Sipsas et al., 1997; Shiga et al., 1996
B*3501 (B35)	RT	175–183	NPDIVIYQY	Sipsas et al., 1997; Shiga et al., 1996
B*3501 (B35)	RT	175–183	HPDIVIYQY	Rowland-Jones et al., 1995
B*3501 (B35)	gp160	42–52	VPVWKEATTTL	Wilkes & Ruhl, 1999
B*3501 (B35)	gp160	78–86	DPNPQEVVL	Shiga et al., 1996
B*3501 (B35)	gp160	606–614	TAVPWNASW	Johnson et al., 1994
B*3501 (B35)	Nef	74–81	VPLRPMTY	Culmann <i>et al.</i> , 1991; Culmann-Penciolelli <i>et al.</i> , 1994

 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
B*3701 (B37)			2 C D F E M	Falk et al., 1993
			L	
B*3701 (B37)	Nef	120–128	YFPDWQNYT	Culmann et al., 1991; Culmann, 1999
B*3801 (B38) B*3801 (B38)	Vif gp160	79–87 104–112	WHLGQGVSI MHEDIISLW	Sabbaj <i>et al.</i> , 2004 Cao, 2002
B*3901 (B39)			2 C <b>R</b> L	Falk <i>et al.</i> , 1995a
B*3901 (B39)	p24	61–69	<b>H</b> GHQAAMQML	Kurane & West, 1999
B*3910 (B39)	p24	48–56	TPQDLNTML	Honeyborne & Kiepiela, 2005
B*4001 (B60)			2 C <b>E L</b>	Falk et al., 1995b
B*4001 (B60) B*4002 (B61)	p17 p24 p2p7p1p6 RT RT gp160 Nef Nef P17 p24 p24 p2p7p1p6 Nef Integrase Integrase	92–101 44–52 118–126 5–12 202–210 805–814 37–45 92–100 11–19 70–78 78–86 64–71 92–100 28–36 260–268 48–56	IEIKDTKEAL SEGATPQDL KELYPLTSL IETVPVKL IEELRQHLL QELKNSAVSL LEKHGAITS KEKGGLEGL GELDRWEKI KETINEEAA AEWDRVHPV TERQANFL KEKGGLEGL LPPIVAKEI VPRRKAKII TPQDLNTML	Altfeld et al., 2000 Altfeld et al., 2000 Yu et al., 2002b Draenert, 2004 Altfeld et al., 2000 Altfeld et al., 2000 Draenert, 2004 Altfeld et al., 2000 Sabbaj et al., 2000 Sabbaj et al., 2003
B*4201 (B42) B*4201 (B42) B*4201 (B42) B*4201 (B42)	p24 RT Nef Nef	48–56 271–279 71–79 128–137	YPGIKVRQL RPQVPLRPM TPGPGVRYPL	Wilkes & Ruhl, 1999 Honeyborne, 2006 Goulder, 1999
B44 B44	Protease gp160	34–42 31–39	EEMNLPGRW AENLWVTVY	Rodriguez <i>et al.</i> , 2004 Borrow <i>et al.</i> , 1997b
B*4402 (B44)			2 C E F Y	Rammensee et al., 1999
B*4402 (B44) B*4402 (B44) B*4402 (B44)	p24 p24 gp160	162–172 174–184 31–40	RDYVDRFYKTL AEQASQDVKNW AENLWVTVYY	Ogg <i>et al.</i> , 1998 Lewinsohn, 1999 Borrow <i>et al.</i> , 1997a

 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
B*4403 (B44)	p17	78–86	LYNTVATLY	Masemola et al., 2004b
B*4415 (B12)	p24	28–36	EEKAFSPEV	Bird et al., 2002
B*4501 (B45)	p2p7p1p6	1–10	AEAMSQVTNS	Sabbaj <i>et al.</i> , 2004
B50	Nef	37–45	LEKHGAITS	Draenert, 2004
B51 B51	Vif	57–66 29–37	IPLGDAKLII EAVRHFPRI	Bansal <i>et al.</i> , 2004 Cao <i>et al.</i> , 2003
B*5101 (B51)	Vpr	29-31	2 C A F P I	Falk <i>et al.</i> , 1995a
B*5101 (B51) B*5101 (B51) B*5101 (B51)	RT RT gp160	42–50 128–135 416–424	<b>G</b> EKEGKISKI TAFTIPSI LPCRIKQII	Haas <i>et al.</i> , 1998; Haas, 1999 Sipsas <i>et al.</i> , 1997 Tomiyama <i>et al.</i> , 1999
B*5201 (B52)			2 C I V	Rammensee et al., 1999
B*5201 (B52)	p24	143–150	Q RMYSPTSI	Wilkes & Ruhl, 1999; Wilson et al., 1997
B53	Nef	135–143	YPLTFGWCF	Kiepiela & Goulder, 2002
B*5301 (B53)			2 C <b>P L</b>	Hill et al., 1992
B*5301 (B53) B*5301 (B53) B*5301 (B53)	p24 p24 Tat	48–56 176–184 2–11	TPYDINQML QASQEVKNW EPVDPRLEPW	Gotch <i>et al.</i> , 1993 Buseyne <i>et al.</i> , 1996, 1997; Buseyne, 1999 Addo <i>et al.</i> , 2001
B*5301 (B53)	Nef	135–143	YPLTFGWCY	Sabbaj et al., 2003
B*5501 (B55)			2 C <b>P</b>	Barber et al., 1995
B*5501 (B55)	gp160	42–51	A VPVWKEATTT	Shankar <i>et al.</i> , 1996; Lieberman, 1999
B57 B57 B57 B57 B57 B57	p24 Protease Integrase Nef Nef	32–40 70–77 123–132 116–124 127–135 137–145	FSPEVIPMF KAIGTVLV STTVKAACWW HTQGYFPDW YTPGPGIRY LTFGWCFKL	Frahm <i>et al.</i> , 2005 Frahm <i>et al.</i> , 2005 Rodriguez <i>et al.</i> , 2004; Addo & Rathod, 2004 Draenert, 2002 Frahm <i>et al.</i> , 2005 Frahm <i>et al.</i> , 2005

 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
B*5701 (B57)			12 C	Barber <i>et al.</i> , 1997
			A F	
			T W	
			S	
			K Y	
B*5701 (B57)	p24	15–23	ISPRTLNAW	Johnson et al., 1991; Goulder et al., 1996b
B*5701 (B57)	p24	30–40	KAFSPEVIPMF	Goulder et al., 1996b
B*5701 (B57)	p24	108–118	TSTLQEQIGWF	Goulder et al., 1996b
B*5701 (B57)	p24	176–184	QASQEVKNW	Goulder et al., 1996b
B*5701 (B57)	RT	244–252	IVLPEKDSW	van der Burg et al., 1997; Hay, 1999
B*5701 (B57)	Integrase	173–181	KTAVQMAVF	Goulder et al., 1996b; Hay, 1999
B*5701 (B57)	Vpr	30–38	AVRHFPRIW	Altfeld et al., 2001a
B*5701 (B57)	Vif	31–39	ISKKAKGWF	Altfeld et al., 2001a
B*5701 (B57)	Rev	14–23	KAVRLIKFLY	Addo et al., 2001
B*5701 (B57)	Nef	116–125	HTQGYFPDWQ	Culmann <i>et al.</i> , 1991
B*5701 (B57)	Nef	120–128	YFPDWQNYT	Culmann et al., 1991
B*5703 (B57)	p24	30-37	KAFSPEVI	Goulder et al., 2000b
B*5703 (B57)	p24	30–40	KAFSPEVIPMF	Goulder et al., 2000b
B58	p17	76–86	RSLYNTVATLY	Frahm et al., 2005
B58	Tat	2-11	EPVDPRLEPW	Frahm & Brander, 2005
B58	gp160	59–69	KAYETEVHNVW	Rathod & Bishop, 2006
B*5801 (B58)			12 C	Barber et al., 1997; Falk et al., 1995b
( 1 1 )			A F	,,,,,
			T W	
			S	
			K	
			V	
			I	
B*5801 (B58)	p24	108-117	TSTVEEQQIW	Bertoletti et al., 1998
B*5801 (B58)	p24	108-117	TSTLQEQIGW	Goulder et al., 1996b
B*5801 (B58)	RT	375–383	IAMESIVIW	Kiepiela & Goulder, 2002
B*5801 (B58)	Rev	14–23	KAVRLIKFLY	Addo et al., 2001
B62	Nef	19–27	RMRRAEPAA	Cao, 2002
B63	p17	76–86	RSLYNTVATLY	Frahm et al., 2005
B63	p24	15-23	ISPRTLNAW	Frahm et al., 2005
B63	p24	30-40	KAFSPEVIPMF	Frahm et al., 2005
B63	Rev	14-23	KAVRLIKFLY	Frahm et al., 2005
B63	Nef	127-135	YTPGPGIRY	Frahm et al., 2005
B63	Nef	137–145	LTFGWCFKL	Frahm et al., 2005
B81	Protease	80–90	TPVNIIGRNML	Honeyborne et al., 2006
B81	RT-Integrase	560-8	LFLDGIDKA	Addo, 2002
B*8101 (B81)	p24	48–56	TPQDLNTML	Goulder et al., 2000a
B*8101 (B81)	Vpr	34–42	FPRIWLHGL	Altfeld et al., 2001a

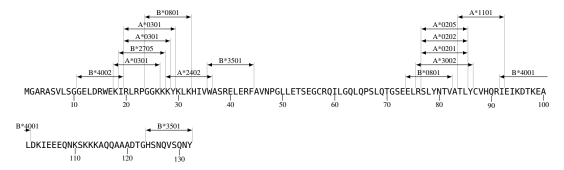
 Table I-A.1: Best defined HIV CTL epitopes (cont.).

HLA	Protein	AA	Sequence	Reference
Cw*0102 (Cw1)			23 C <b>A</b> L L P	Barber <i>et al.</i> , 1997
Cw*0102 (Cw1) Cw*0102 (Cw1)	p24 Gag-Pol TF	36–43 24–31	VIPMFSAL NSPTRREL	Goulder <i>et al.</i> , 1997a Liu <i>et al.</i> , 2006
Cw3	Nef	83–91	AALDLSHFL	Draenert, 2004
Cw*0303 (Cw9)	p24	164–172	YVDRFFKTL	Honeyborne, 2003
Cw*0304 (Cw10) Cw*0304 (Cw10)	p24 gp160	164–172 557–565	YVDRFFKTL RAIEAQQHL	Honeyborne, 2003 Currier <i>et al.</i> , 2002; Trocha, 2002
Cw*0401 (Cw4)			2 6 C Y L P F F M	Falk et al., 1994
Cw*0401 (Cw4)	gp160	375–383	SFNCGGEFF	Wilson et al., 1997; Johnson et al., 1993
Cw5	p24	174–185	AEQASQEVKNWM	Draenert, 2004
Cw*0501	Rev	67-75	SAEPVPLQL	Addo et al., 2001
Cw6	Nef	120-128	YFPDWQNYT	Frahm & Brander, 2005
Cw7 Cw7	Nef Nef	105–115 105–115	KRQEILDLWVY RRQDILDLWIY	Kiepiela & Goulder, 2002 Yu <i>et al.</i> , 2002a
Cw8 Cw8	gp160 Nef	557–565 82–91	RAIEAQQHM KAAVDLSHFL	Bishop & Honeyborne, 2006 Harrer <i>et al.</i> , 1996c
Cw*0802 (Cw8)	p24	48–56	TPQDLNTML	Goulder <i>et al.</i> , 2000a; Honeyborne & Kiepiela, 2005
Cw*0802 (Cw8) Cw*0802 (Cw8) Cw*0802 (Cw8)	RT Nef Nef	495–503 83–91 83–91	IVTDSQYAL AAVDLSHFL GAFDLSFFL	Rathod & Honeyborne, 2006 Cao <i>et al.</i> , 2003 Rathod & Honeyborne, 2006
Cw*0804 (Cw8)	p17	33–41	HLVWASREL	Masemola et al., 2004b
Cw12	Tat	30–37	CCFHCQVC	Cao et al., 2003; Nixon et al., 1999
Cw14	p17	78–85	LYNTVATL	Horton & Havenar-Daughton, 2005
Cw15	gp160	557–565	RAIEAQQHL	Trocha, 2002
Cw18 Cw18 Cw18 Cw18	p24 p24 Integrase gp160	142–150 161–169 165–172 511–519	VRMYSPVSI FRDYVDRFF VRDQAEHL YRLGVGALI	Honeyborne, 2006 Honeyborne & Kiepiela, 2005 Rathod & Honeyborne, 2006 Honeyborne, 2006

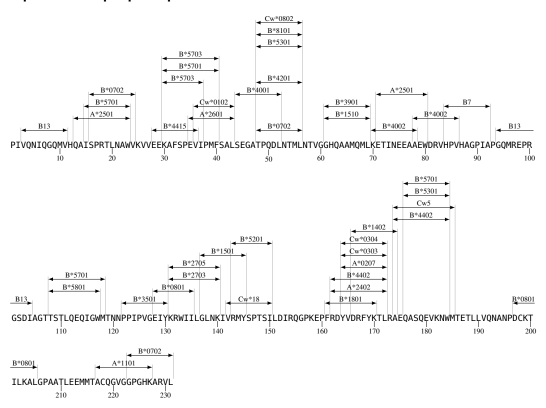
#### I-A-5 Map of optimal HIV-1 CTL epitopes

The location and HLA restriction elements of CTL epitopes are indicated on protein sequences of HXB2. These maps are meant to provide the relative location of defined epitopes on a given protein, but the HXB2 sequence may not actually carry the epitope of interest, as it may vary relative to the sequence for which the epitope was defined.

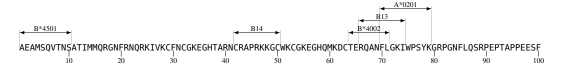
#### p17 Optimal CTL Epitope Map

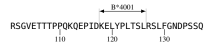


#### p24 Optimal CTL Epitope Map



#### p2p7p1p6 Optimal CTL Epitope Map





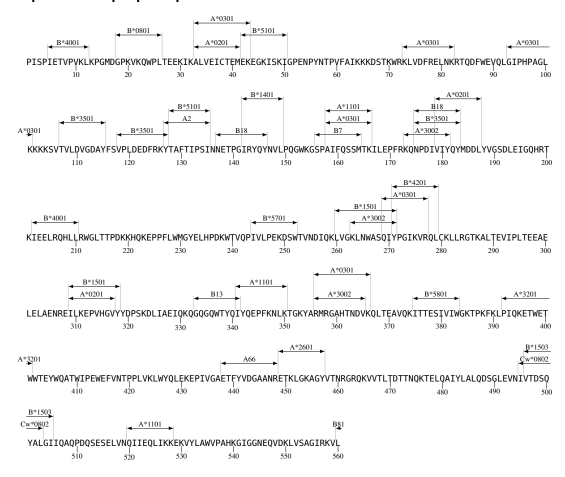
#### **Gag/Pol TF Optimal CTL Epitope Map**



#### **Protease Optimal CTL Epitope Map**

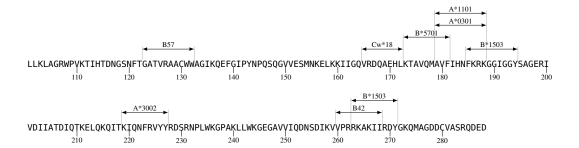


#### **RT Optimal CTL Epitope Map**

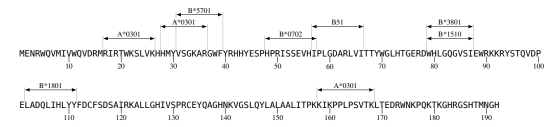


#### **Integrase Optimal CTL Epitope Map**

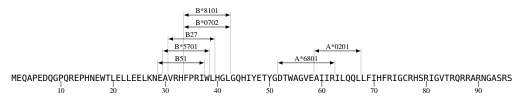




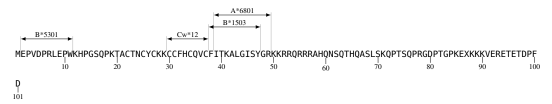
#### Vif Optimal CTL Epitope Map



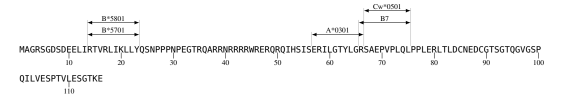
#### **Vpr Optimal CTL Epitope Map**



#### **Tat Optimal CTL Epitope Map**



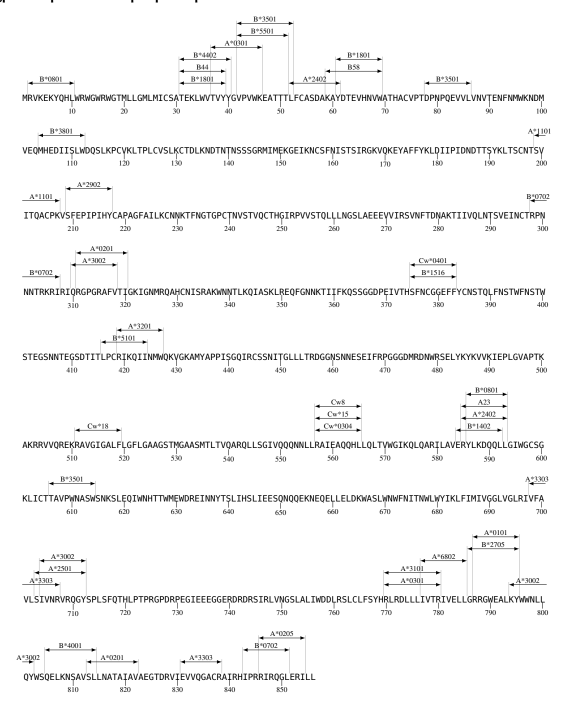
#### **Rev Optimal CTL Epitope Map**



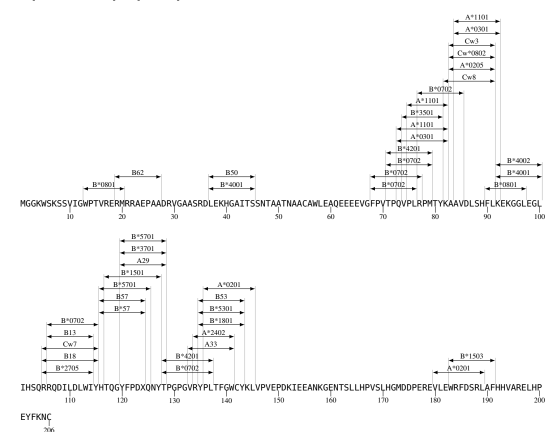
#### **Vpu Optimal CTL Epitope Map**



#### gp160 Optimal CTL Epitope Map



#### **Nef Optimal CTL Epitope Map**



#### **I-A-6** Acknowledgments

We would like to express our gratitude to those researchers in the field who continuously contribute to this database. The mostly unpublished data added to this year's update stemming from the AIDS Research Center at Mass. General Hospital were largely funded by two NIH contracts (NO1-A1-15442, NO1-A1-30024) supporting HLA typing and HIV CTL epitope definition in non-Caucasian populations and non-clade-B HIV infection as well as R01-A1-067077 assessing the promiscuous presentation of HLA class I restricted epitopes.

We very much welcome any criticism, comments and additions to this list since we are sure that some epitopes will unintentionally escape our attention, despite close monitoring of the literature. Please write or call us with any comments you may have.

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### Beyond *Mamu-A\*01* + Indian Rhesus Macaques: Continued Discovery of New MHC Class I Molecules that Bind Epitopes from the Simian AIDS Viruses

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#### I-B-1 Introduction

Developing an effective vaccine against HIV is a global public health priority. Nonhuman primate research will play a critical role in such development as simian immunodeficiency virus (SIV) infection of nonhuman primates is the best animal model available for AIDS, and nonhuman primates are used for testing of vaccination strategies. In the past decade, substantial progress has been made in defining and characterizing T cell responses for this animal model.

Accumulating evidence suggests that cellular immune responses play a major role in controlling HIV and SIV replication, directing recent efforts to vaccine regimens that elicit CD8+ T lymphocyte responses [McMichael & Rowland-Jones, 2001]. A variety of studies have shown associations between certain MHC class I alleles in both slow and rapid HIV/SIV disease progression [Carrington & O'Brien, 2003; Bontrop & Watkins, 2005]. Of particular interest are "elite controllers" (ECs), rare individuals who spontaneously control HIV/SIV viremia to very low levels (<50 vRNA copies/ml of plasma for HIV-infected humans [Pereyra *et al.*, 2007], <1,000 vRNA copies/ml of plasma for SIV-infected macaques [Yant *et al.*, 2006; Loffredo *et al.*, 2007b]). Understanding this natural control may aid in the development of an effective HIV vaccine.

There are, unfortunately, many difficulties inherent to studying HIV-infected humans. Viral control appears to be mediated during resolution of acute phase viremia, with the appearance of CD8+ T cell responses in both HIV-infected humans and SIV-infected macaques [Borrow *et al.*, 1994; Koup *et al.*, 1994; Reimann *et al.*, 1994; Yasutomi *et al.*, 1993; Kuroda *et al.*, 1999]. HIV is rarely diagnosed during acute infection however [Weintrob *et al.*, 2003; Kuo *et al.*, 2005; Mayben *et al.*, 2007], making the study of immune responses involved in initial control of HIV replication extremely difficult. This is further complicated by the diversity of HIV isolates with which individuals might be infected.

AIDS research with nonhuman primates provides an animal model to complement human studies. Nonhuman primates can provide examples of successful immune containment of pathogenic immunodeficiency virus replication, and researchers have direct control over key variables such as virus strain, host genotype, and route of infection. Perhaps most importantly, the immunology and pathogenesis of acute infection can easily be studied in macaques. Overall, studying immune responses to SIV in macaques is simpler than studying immune responses to HIV in humans. All macaques can be infected with a known, often clonal, viral stock. Infection with a defined viral stock enables complete and accurate immunological tracking of early immune responses by the use of corresponding peptides in ex vivo immunological assays. Moreover, the timing of immune responses after infection, the associated viral sequence evolution, and plasma virus concentrations may be closely monitored.

Most published studies of CD8+ T cell responses in Indian rhesus macaques have used animals which express the common MHC class I allele *Mamu-A\*01*. This allele was found to be associated with lower set-point viremia in vaccinated and non-vaccinated macaques in several studies [Mao *et al.*, 2005; Mühl *et al.*, 2002; Pal *et al.*, 2002; Zhang *et al.*, 2002]. Such animals have been used largely because their CD8+ T cell responses were the first to be exhaustively identified, and *Mamu-A\*01* can be readily identified using sequence-specific PCR primers [Knapp *et al.*, 1997]. The Mamu-A\*01 peptide binding motif has been defined, and a comprehensive scan of the SIVmac239

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In *HIV Molecular Immunology 2006/2007*. Bette T. M. Korber, Christian Brander, Barton F. Haynes, Richard Koup, John P. Moore, Bruce D. Walker, and David I. Watkins, editors. Publisher: Los Alamos National Laboratory, Theoretical Biology and Biophysics, Los Alamos, New Mexico. p. 29–51.

proteome yielded fourteen epitopes restricted by this MHC class I molecule [Allen *et al.*, 1998, 2001]. Tetramers are available for several of these epitopes, facilitating detailed vaccine and pathogenesis studies [Kuroda *et al.*, 1998; Allen *et al.*, 2001; Mothé *et al.*, 2002a; Egan *et al.*, 1999; Allen *et al.*, 2000].

As described in the previous compendium update and elsewhere, this focus on Mamu-A\*01+ macaques has created a bottleneck in AIDS vaccine research due to a shortage of animals expressing this allele [O'Connor et al., 2002b; Cohen, 2000]. Compared to the human system, only a small fraction of macaque MHC class I alleles have been characterized at this time [Bontrop & Watkins, 2005]. Since the publication of the last SIV compendium review in 2001 [O'Connor et al., 2002b], significant advances have been made in understanding nonhuman primate immunogenetics. Two MHC class I alleles, Mamu-B\*17 and Mamu-B\*08, were found to be enriched in EC cohorts and associated with reduced SIV replication in the chronic phase of infection [Yant et al., 2006; Loffredo et al., 2007b]. The binding motifs of four additional Indian rhesus macaque MHC class I alleles have been defined, and the SIVmac239 epitopes presented by these alleles identified [Mothé et al., 2002b; Loffredo et al., 2004; Sette et al., 2005; Loffredo et al., 2005]. Knowledge of the minimal-optimal epitopes restricted by these alleles should allow the use of many more Indian rhesus macaques in studies of CD8+ T cell dynamics following SIV infection, as well as a fuller understanding of the immune responses generated against SIV. Additionally, several other species of macaques are now being used in AIDS research, and their cellular immune responses are beginning to be defined.

This review highlights recent advances in AIDS research using nonhuman primate models and provides a comprehensive list of the published MHC class I and class II epitopes (Table I-B.1 and Table I-B.2). A summary of the peptide-binding motifs of several macaque MHC alleles is also presented (Table I-B.3).

# I-B-2 Indian rhesus macaques: MHC class I molecules and CD8+ T cell epitopes

The Indian rhesus macaque (*Macaca mulatta*) remains the best characterized and most commonly studied nonhuman primate AIDS model. At this time, the peptide binding motifs of five MHC class I molecules (Mamu-A\*01, -A\*02, -A\*11, -B\*01, and -B\*17) have been fully defined in the SIV-infected rhesus macaque model (Table I-B.3), enabling the identification of 43 of the 66 known SIV-specific CD8+ T cell epitopes (Table I-B.1) [Allen *et al.*, 1998, 2001; Mothé *et al.*, 2002b; Loffredo *et al.*, 2004; Sette *et al.*, 2005; Loffredo *et al.*, 2005]. As described

above, Mamu-A\*01 has been intensely investigated, as it was the first MHC class I molecule studied in detail [Allen et al., 1998; Furchner et al., 1999; Allen et al., 2001, 2000; Egan et al., 1999; Barouch et al., 2002; Mothé et al., 2002a; O'Connor et al., 2002a; Zhang et al., 2002; Pal et al., 2002; Mühl et al., 2002; O'Connor et al., 2004; Friedrich et al., 2004b; Knapp et al., 1997; Peyerl et al., 2003; Kuroda et al., 1998; Loffredo et al., 2007a]. The addition of four well-characterized MHC class I molecules significantly broadens the repertoire of immune responses that researchers may effectively monitor in animal studies. Mamu-A\*02 and Mamu-B\*17 are particularly helpful with twenty [Vogel et al., 2002a; Loffredo et al., 2004; Robinson et al., 2001; Watanabe et al., 1994] and twelve [Evans et al., 1999, 2000; Dzuris et al., 2000; Mothé et al., 2002b; Horton et al., 2001] epitopes, respectively, known to be restricted by these two alleles. In addition, each is present at a high frequency (>10%), in most captive rhesus macaque colonies [Kaizu et al., 2007].

Mamu-B\*01, another high frequency allele, is unusual in that it does not appear to be involved in the immune response directed against SIVmac239 [Loffredo et al., 2005], although previous published reports had described six Mamu-B\*01-restricted CD8+ T cell epitopes [Yasutomi et al., 1995; Su et al., 2005]. However, the MHC restriction of these six putative Mamu-B\*01-restricted epitopes was not verified with MHC class I transfection of 721.221 cells [Yasutomi et al., 1995; Su et al., 2005]. In a subsequent study, it was found that the putative epitopes did not bind to Mamu-B\*01 with biologically relevant affinities [Loffredo et al., 2005], nor were CD8+ responses to these peptides detected in eight Mamu-B\*01+ SIV-infected Indian rhesus macaques. Hence, vaccine immunogenicity studies in Mamu-B\*01+ macaques may not yield useful results.

## I-B-3 Indian rhesus macaques: web-based immunogenetics resources

Based on the identified SIV-specific CD8+ T cell epitopes and the peptide binding profiles of their corresponding Indian rhesus macaque MHC class I alleles, a computational algorithm was developed to predict the peptide binding and potential T cell epitopes restricted by the MHC class I molecules Mamu-A\*01, -A\*02, -A\*11, -B\*01, and -B\*17 [Peters et al., 2005a]. This can be accessed at the website http://www.mamu.liai.org/. Previous studies defined detailed peptide binding motifs, aiding discovery of the majority of identified SIVmac239 epitopes [Allen et al., 1998, 2001; Mothé et al., 2002b; Loffredo et al., 2004; Sette et al., 2005; Loffredo et al., 2005]. However, it is possible that epitope screens done in chronically infected macaques failed to detect some subdominant responses restricted by these alleles. Identification and characterization

of additional subdominant responses may be aided by using the website's search engine to find the predictive binding values of peptides within regions of novel responses of unknown restriction. In addition, this resource may be of use to researchers studying other viral pathogens in the Indian rhesus macaque, expediting CD8+ T cell epitope identification. The Immune Epitope Database and Analysis Resource (IEDB) [Peters et al., 2005b], located at http://www.immuneepitope.org/, is another useful resource recently made available. The IEDB project is hosted at the La Jolla Institute for Allergy and Immunology (LIAI), and one of its foci is the compilation of known immune (antibody and T cell) epitopes and MHC binding data. These two resources provide useful information regarding the epitopes and alleles discussed in this review, in addition to covering humans, rodents, and various other animal species.

## I-B-4 Indian rhesus macaques: viral evolution and epitope cross-reactivity

As more epitopes restricted by different Indian rhesus macaque alleles have been identified and followed over the course of SIV infection, sequencing of viral RNA has revealed selection of variation by CD8+ T cells. Escape mutations are detected in many epitopes targeted by CD8+ T cells, and variation is found in epitopes restricted by each of the alleles discussed previously in the review [Evans et al., 1999; Allen et al., 2000; O'Connor et al., 2002a, 2004; Barouch et al., 2002, 2003; Peyerl et al., 2003; Friedrich et al., 2004b; Loffredo et al., 2004; Vogel et al., 2002a]. This variation is selected for with varying kinetics following infection, though for a given epitope the same escape mutation is often found in multiple animals. Recent studies have also shown that viral escape from CD8+ T cell pressures can exact a fitness cost to the virus via fitness assays and reversion experiments [Friedrich et al., 2004a; Peyerl et al., 2003; Friedrich et al., 2004b; Barouch et al., 2005].

Reactivity to escape variant peptides is frequently detectable in enzyme-linked immunospot or intracellular cytokine staining assays. However, this putative "cross-reactivity" may be an artifact of the non-physiological antigen presentation that occurs when high concentrations of exogenous antigen are used to stimulate T cells. For at least two SIVmac239 epitopes, Mamu-A\*01-restricted Gag<sub>181-189</sub>CM9 and Tat<sub>28-35</sub>SL8, consistent reactivity to escape variant peptides has been observed, albeit with reduced functional avidity. By contrast, cells infected with corresponding escape variant viruses were not recognized by CD8+ T cell lines specific for the wild-type epitope [Loffredo *et al.*, 2007a; Valentine *et al.*, 2007]. Whether *de novo* responses specific for escape variant sequences

are generated has not been rigorously studied.

## I-B-5 Indian rhesus macaques: natural containment of pathogenic SIV replication

The most interesting, recent studies of Indian rhesus macaque immunogenetics involved the characterization of two MHC class I alleles associated with slow disease progression in SIV-infected macaques. Both Mamu-B\*17 and Mamu-B\*08, unlike Mamu-A\*01, are MHC class I alleles significantly enriched in elite controller (EC) cohorts [Yant et al., 2006; Loffredo et al., 2007b]. Fourteen of the sixteen (88%) EC macaques identified from a cohort of 196 SIVmac239-infected macaques expressed either Mamu-B\*17 or Mamu-B\*08. This percentage is similar to a previous study that demonstrated the overrepresentation of HLA-B\*5701 (11 of 13, 85%) in a cohort of long term nonprogressors / ECs with normal CD4+ T cells counts and HIV replication less than 50 vRNA copies/ml [Migueles et al., 2000]. Mamu-B\*17+ and Mamu-B\*08+ macaques also exhibit a greater reduction in SIV viremia compared to Mamu-A\*01+ macaques in a cohort of 196 SIVmac239-infected macaques [Yant et al., 2006; Loffredo et al., 2007b].

Intriguingly, the peptide binding motif of Mamu-B\*17 is broadly similar to that of HLA-B57, while the Mamu-B\*08 motif resembles that of HLA-B27. HLA-B57 and -B27 are associated with lower plasma virus concentrations in HIV-infected individuals [Carrington & O'Brien, 2003]. Although Mamu-B\*17 seems to tolerate a wider array of residues at position two than HLA-B57, both molecules require W, F, or Y at the C-terminal anchor position [Mothé et al., 2002b; Rammensee et al., 1999; Marsh et al., 2000]. A previous study demonstrated that another macaque MHC class I molecule, Mamu-B\*03, binds peptides conforming to the HLA-B27 binding motif [Dzuris et al., 2000]. This allele was also associated with slow SIV disease progression [Evans et al., 1999], but additional studies were hindered by the low frequency of this allele in captive rhesus macaques (overall < 1%; Kaizu et al. [2007]). Mamu-B\*03 and Mamu-B\*08 are almost identical in amino acid sequence [Boyson et al., 1996], with only two amino acid differences between Mamu-B\*03 and Mamu-B\*08 in regions that influence peptide binding and antigen recognition [Bjorkman et al., 1987; Garrett et al., 1989]. Both differences reside in the alpha-1 domain (exon two). Therefore, it is likely that Mamu-B\*08 also shares this HLA-B27 binding profile. This hypothesis is supported by the fact that all seven Mamu-B\*08-restricted CD8+ T cell epitopes currently described (Table I-B.1) fit the peptide-binding motif for HLA-B27 of an R at position two and an L at the C-terminus [Loffredo et al., 2007c]. Studies to completely define the Mamu-B\*08 peptide binding motif are ongoing, and should help to identify the complete repertoire of SIVmac239-specific CD8+ T cell epitopes restricted by Mamu-B\*08.

The discovery of Mamu-B\*17 and Mamu-B\*08 now provide us with a unique opportunity not previously available due to technological and assay limitations. Currently, we have found that about 20–25% of *Mamu-B\*17+* and about 50% of *Mamu-B\*08+* SIV-infected Indian rhesus macaques are elite controllers [Yant *et al.*, 2006; Loffredo *et al.*, 2007b]. With peptide binding motifs similar to the analogous human EC alleles *HLA-B57* and *HLA-B27*, respectively, and definition of at least seven SIV epitopes for each macaque allele (Table I-B.1), investigators now have the resources to study successful immune responses against AIDS viruses in the SIV-infected macaque model.

## I-B-6 Indian rhesus macaques: MHC class II molecules and CD4+ T cell epitopes

As with HIV, mapping of CD4+ T cell epitopes in SIV has lagged behind studies of CD8+ T cell epitopes. This is due in part to the difficulty in maintaining such responses in the face of ongoing viral replication, as HIV-specific CD4+ T cells are preferentially infected [Douek et al., 2002]. Initially, studies focused on identifying Env-specific CD4+ T cell responses in SHIV-infected macaques [Lekutis & Letvin, 1997; Dzuris et al., 2001; Lekutis et al., 1997]. Two MHC class II molecules, Mamu-DRB1\*0406 and Mamu-DRB\*w201, presented three Env-specific CD4+ T cell responses, enabling an Env-specific MHC class II tetramer to be constructed [Kuroda et al., 2000]. Later, peptide binding motifs for these two alleles were generated, and used to identify other SIV-derived peptides that Mamu-DRB1\*0406 and Mamu-DRB\*w201 might present [Dzuris et al., 2001]. This investigation also discussed two new novel CD4+ T cell responses against Gag and Rev that were restricted by Mamu-DRB\*w201. Unpublished data from Giraldo Vela et al. describe five additional MHC class II alleles that restrict CD4+ T cell responses in Gag, Rev, Nef, and Vpx, thereby doubling the number of known SIV/SHIV epitopes (Table I-B.2). Numerous other regions of SIV have been documented to contain CD4+ T cell epitopes [Mills et al., 1991; Sarkar et al., 2002; Vogel et al., 2002b]. However, the MHC class II restriction of the majority of these responses remains undefined.

## I-B-7 Alternatives to the SIV-infected Indian rhesus macaque animal model

The demand for Indian rhesus macaques has lessened somewhat with a growing interest in alternative nonhuman

primate models for AIDS. Currently, pigtail macaques (*Macaca nemestrina*) are being used in epitope identification and vaccine/pathogenesis studies. In addition, vaccine-induced immune responses have been effective at controlling SIVmac239 replication in Burmese macaques (*Macaca mulatta*), while the simple MHC genetics of Mauritian cynomolgus macaques (*Macaca fascicularis*) may offer unique opportunities for future SIV studies. There are, however, some key differences between these groups of animals, which are pertinent to SIV research.

All of these macaques, of the genus Macaca, are native to Asia and are non-natural hosts for SIV. Burmese-origin rhesus macaques, while the same species as Indian-origin rhesus macaques, are from a geographically separate population and possess divergent MHC class I alleles. Different subpopulations of rhesus macaques (Indian, Burmese, and Chinese origin) can be readily distinguished by mitochondrial DNA sequencing [Smith & McDonough, 2005], and by single nucleotide polymorphism analysis [Ferguson et al., 2007; Malhi et al., 2007], tools that may be useful for determining ancestry of animals with unknown origin. The most recent common ancestor of the rhesus macaque species is estimated to have lived about 1.9 million years ago, around the time rhesus macaques diverged from cynomolgus macaques [Hernandez et al., 2007]. Pigtailed macagues are somewhat more distantly related, having separated from the rhesus macaque lineage about 3.5 million years ago [Morales & Melnick, 1998].

These different groups of macaques experience varying degrees of pathogenicity following infection with commonly used SIV isolates. A comparative study by Reimann et al. [2005] found no significant difference in the acutephase peak of viremia in SIVmac251-infected Indian rhesus, Chinese rhesus, and cynomolgus macaques, while a separate study of SIVmac251-infected pigtail macaques measured a very similar acute-phase viral peak [Batten et al., 2006]. Differences in plasma virus concentrations emerge after acute infection, the timing of which suggests a determinative role for the adaptive immune response, rather than an inherent difference in the replicative capacity of SIV in these animals. Overall, rhesus macaques of Indian descent consistently have the highest viral set-points. In comparison, it appears that the chronicphase plasma viremia of SIVmac239 in Burmese rhesus macaques (geometric mean of 65,000 vRNA copies/ml in one small study [Yamamoto et al., 2007]) is similar, or perhaps slightly lower than that observed in a large cohort of Indian rhesus macaques (geometric mean 223,800 vRNA copies/ml, [Loffredo et al., 2007b]). However, a direct comparison of these animals has not been published. Meanwhile, cynomolgus macaques have significantly reduced plasma virus concentrations (approximately twologs) compared to Indian-origin rhesus macaques, with a large proportion of these cynomolgus macaques becoming ECs after SIVmac251 challenge [Reimann et al., 2005]. Pigtail macaques are also productively infected with SIV-mac251, with substantial variability in their viral set-point having been documented [Batten *et al.*, 2006].

Why different macaque species/subspecies have different disease courses following infection with the same virus has not been defined, but it is notable that SIV isolates used in the majority of studies (SIVmac239, SIVmac251, and SHIV-89.6P) have been passaged in rhesus macaques of Indian origin. One study has shown that passaging SIV in rhesus macaques of Chinese origin resulted in increased viral loads in subsequently infected Chinese macaques [Burdo *et al.*, 2005]. While the mutations acquired were not characterized, this study suggests that species-specific adaptations might play a role in determining viral set-point for different groups of animals.

#### I-B-8 Burmese rhesus macaques

Several interesting vaccine experiments have utilized SIVinfected Burmese rhesus macaques. A Gag-expressing DNA-prime/Sendai virus vector boost vaccination led to control of SIVmac239 replication in five of eight vaccinated Burmese macaques [Matano et al., 2004]. By week five, the successful vaccinees had undetectable plasma virus concentrations. Interestingly, viruses from these five animals had escape mutations in Gag that appeared to have occurred at a high fitness cost to the virus [Matano et al., 2004; Kobayashi et al., 2005]. The escape analysis demonstrated that three vaccinees shared an MHC class I haplotype (90-120-Ia), which restricted some of the Gag-specific CD8+ T cell responses (Table I-B.1). Follow-up studies in these animals have shown that multiple Gag-specific CD8+ T lymphocyte responses were involved in the vaccine-induced control [Kawada et al., 2006], and additional viral mutations associated with this MHC class I haplotype (90-120-Ia) accumulated. At this time, responses have not been mapped to specific MHC class I alleles.

Additionally, SIV-specific CD4 responses in the Burmese macaques are being investigated and appear to be associated with viral control [Lun *et al.*, 2004]. SIV-specific CD8+ T cells that are not directed against Gag also appear to be important in Burmese macaques, although responses have not yet been mapped [Kawada *et al.*, 2007].

#### **I-B-9** Pigtail macagues

Until recently, research with pigtail macaques had been hampered by a lack of defined viral epitopes and limited characterization of pigtail macaque MHC class I alleles. However, over the past several years considerable effort has been directed toward understanding the immunogenetics of pigtail macaques, widening the resources available to researchers. Importantly, many *Macaca nemestrina* MHC

class I alleles (*Mane*) have now been characterized from more than 100 pigtail macaques [Lafont *et al.*, 2003; Pratt *et al.*, 2006; Smith *et al.*, 2005a].

Immunogenetic analyses of these animals have identified several defined minimal-optimal epitopes in Gag (Table I-B.1). Of particular interest is an immunodominant response, Gag<sub>164-172</sub>KP9, restricted by the high frequency allele, Mane-A\*10 (and later Mane-A\*16, an MHC class I allele of similar sequence) [Smith et al., 2005a,b]. This SIV epitope Gag<sub>164-172</sub>KP9 has been studied extensively and parallels can be drawn to the Mamu-A\*01-restricted CD8+ T cell epitope Gag<sub>181-189</sub>CM9. Viral escape at position two of this epitope leads to loss of epitope recognition [Fernandez et al., 2005]. Viral fitness was also impaired by this mutation, as evidenced by sequence reversion when the selective pressure of the Gag<sub>164-172</sub>KP9-specific immune responses was removed [Fernandez et al., 2005; Loh et al., 2007; Fernandez et al., 2007]. In addition, unvaccinated pigtail macaques infected with SIVmac251 that directed a response against Gag<sub>164-172</sub>KP9 have significantly reduced plasma viremia compared to Gag<sub>164-172</sub>KP9 nonresponders [Smith et al., 2005a].

Additional research has also identified three subdominant Gag responses in pigtail macaques restricted by various other MHC class I alleles (Table I-B.1) [Smith *et al.*, 2005a; Fernandez *et al.*, 2005; Pratt *et al.*, 2006]. Furthermore, several other Gag-specific CD8 and CD4 responses are currently being mapped, and their restriction elements determined [Fernandez *et al.*, 2005]. Finally, another study provided a comparative virological and immunological analysis of a variety of primate lentiviruses showing the variable pathogenicity of a variety of SIV/SHIVs that are available for use in pigtail macaques [Batten *et al.*, 2006].

#### I-B-10 Mauritian cynomolgus macaques

Research with cynomolgus macaques has not been as common as research with rhesus macaques because many SIV and SHIV strains are less pathogenic in the cynomolgus species [Reimann et al., 2005]. However, lately there has been renewed interest in studying the SIV-specific immune responses in these animals, largely due to the unique immunogenetics of cynomolgus macaques from the Indian Ocean island of Mauritius. During the definition of 66 MHC class I alleles in cynomolgus macaques of Chinese, Vietnamese, and Mauritian origin, it was discovered that most MHC class I alleles could be divided by geographic origin, with few alleles shared by animals descended from geographically distinct populations [Krebs et al., 2005]. This is similar to findings in Chinese and Indian rhesus macaques, which do not express the same MHC class I alleles [Ling et al., 2002; Trichel et al., 2002]. These data highlight the point that macaques from different origins are not interchangeable in studies of cellular immunity.

However, also of great interest was the finding that more than 50% of Mauritian cynomolgus macaques shared a combination of three MHC class I alleles. In contrast, the most frequent Indian rhesus macaque alleles are present at a frequency of only 25-30% in any given population [Kaizu *et al.*, 2007].

Follow-up analysis using microsatellite markers revealed that the Mauritian cynomolgus macaques have extremely simple MHC genetics, with six distinct chromosomal haplotypes accounting for almost all of the MHC class I and class II diversity in this population, likely due to these animals being recently descended from a small founder population [Lawler et al., 1995]. Remarkably, 39% of Mauritian cynomolgus macaques carry at least one copy of the most frequent MHC haplotype, with 8% being homozygous for this haplotype [Wiseman et al., 2007]. This extensive sharing of MHC haplotypes is unprecedented among macaques [Otting et al., 2005; Penedo et al., 2005; Krebs et al., 2005; Wiseman et al., 2007] and could expand the scope of SIV studies undertaken in nonhuman primates by allowing greater control over genetic variability. Studies are now possible in which entire MHC class I haplotypes are matched, rather than a single MHC class I molecule, as is typical in Indian rhesus macaques studies. This may simplify studying cellular immune responses by eliminating unknown influences of unmatched MHC class I alleles. In addition, adoptive lymphocyte transfers that are possible in inbred mouse strains might now be technically feasible with MHC-identical macaques. Recently, the MHC class II alleles for the six common haplotypes were characterized in these animals, enabling researchers to generate useful molecular reagents for CD4+ T cell studies [O'Connor et al., 2007].

At this point, SIV epitopes restricted by these high frequency MHC class I alleles have not been mapped in Mauritian cynomolgus macaques. However, a recent study defined a single Gag minimal-optimal epitope of unknown MHC class I restriction [Negri *et al.*, 2006]. In addition, a similar breadth of cellular immune responses in Gag, Pol, Tat, Env, Rev, and Nef was identified in two MHC class I-identical SIVmac239-infected macaques. Corresponding viral variation was seen in Tat, Rev, and Nef that may prove to be escape from CD8+ T cell pressures [Wiseman *et al.*, 2007].

## I-B-11 Advances in MHC genotyping methods

To support efficient utilization of limited nonhuman primate resources and maximize the understanding of SIV-specific immune responses in vaccine and viral pathogenesis studies, accurate and efficient MHC genotyping technologies are needed.

PCR amplification with sequence specific primers (PCR-

SSP) is currently the preferred method for MHC genotyping at allelic-level resolution in many human clinical laboratories [Olerup & Zetterquist, 1991, 1992] because this technique is specific, robust, and straightforward. The PCR-SSP platform enables rapid, high throughput analysis, yet it is cost effective and requires inexpensive equipment. These benefits have made it a productive choice for MHC class I genotyping of Indian rhesus macaques. Initial investigations designed PCR-SSP primers for a single MHC class I allele of interest, with unique PCR amplification conditions [Knapp et al., 1997; Vogel et al., 2002a; Horton et al., 2001; Loffredo et al., 2005; Robinson et al., 2001; Schramm et al., 2001; Su et al., 2005]. However, recent technological improvements have allowed the use of unified PCR conditions, allowing simultaneous PCR amplification of eight Indian rhesus macaque MHC class I alleles [Kaizu et al., 2007]. These alleles were selected since they have been implicated in the restriction of SIVspecific CD8+ T cell epitopes. Molecular genotyping of Mamu-A\*01, -A\*02, -A\*08, -A\*11, -B\*01, -B\*03, -B\*04, and -B\*17 can now be conducted in a high throughput fashion, using genomic DNA as a template.

Although the importance of defining the complete MHC class I repertoire of a given macaque is well recognized, development of a simple, comprehensive molecular genotyping method for rhesus macaque MHC class I alleles has been difficult. During the evolution of this species, the *Mamu-A* and *Mamu-B* loci were duplicated [Bontrop & Watkins, 2005; Otting *et al.*, 2005; Daza-Vamenta *et al.*, 2004; Boyson *et al.*, 1996; Kulski *et al.*, 2004]. Therefore, macaques express a variable number of MHC class I alleles per haplotype.

The complexity of the nonhuman primate MHC is problematic for PCR-SSP-based genotyping since this technique can only detect known MHC alleles. Moreover, it is difficult to develop allele-specific primers in the absence of a reasonably complete allele database. The established methods to identify all of the alleles in a given macaque involve PCR cloning and sequencing, or generating macaquederived cDNA libraries. While cDNA libraries are the gold standard in identifying MHC alleles, the process is very time consuming and expensive. PCR cloning and sequencing is a less time consuming alternative and requires less starting material. However, this technique may introduce PCR artifacts or provide incomplete MHC coverage due to primer mismatches. Therefore, alternative genotyping methods for rapid identification of all of the alleles in a given animal are being investigated.

The most developed of these techniques is reference strand-mediated conformational analysis, or RSCA. RSCA is a modified heteroduplex assay capable of characterizing complex gene families [Argüello & Madrigal, 1999; Argüello *et al.*, 2003; Kennedy *et al.*, 2005; Krebs *et al.*, 2005]. Heteroduplexes are created by hybridization between fluorescently labeled reference strands and individ-

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ual MHC alleles. RSCA can resolve individual MHC alleles because each MHC/reference strand heteroduplex produces a characteristic and reproducible mobility profile when electrophoresed in a nondenaturing polyacrylamide gel. The mobility profile of each MHC allele can be further resolved for higher degrees of sensitivity with the addition of alternative reference strands. RSCA, unlike conventional genotyping techniques (i.e. PCR-SSP), can detect multiple known and unknown MHC alleles in a single reaction, enabling the researcher to define the full complement of expressed class I alleles in a given animal.

RSCA has been used in several nonhuman primate studies, including the SIV models summarized in this review. These applications include MHC genotyping, allele discovery, epitope restriction determination, and identifying haplotype inheritance patterns [Krebs et al., 2005; Smith et al., 2005a,b; Pratt et al., 2006; Baquero et al., 2006; Wiseman et al., 2007; Tanaka-Takahashi et al., 2007]. Current RSCA platforms can be easily expanded to incorporate newly discovered alleles once a mobility profile is established. However, RSCA is best utilized in tandem with other MHC genotyping techniques due to limited sensitivity in some circumstances [Pratt et al., 2006; Tanaka-Takahashi et al., 2007]. In addition, this technique is still technically demanding [Krebs et al., 2005; Pratt et al., 2006; Smith et al., 2005a,b], so genotyping via capillary electrophoresis is currently being explored for higher throughput and increased sensitivity.

Another alternative for rapid MHC genotyping is microsatellite analysis, also known as short tandem repeat (STR) genotyping. For years, microsatellite genotyping has been applied for tissue matching and donor screening in human transplantation [Carrington & Wade, 1996; Foissac et al., 2001]. MHC genotyping using microsatellite markers has become more common because the method is sensitive, accurate, and cost effective. With the recent publication of the genomic sequence of the entire rhesus macaque MHC [Daza-Vamenta et al., 2004], and more recently the complete genome of an Indian-origin rhesus macaque [Gibbs et al., 2007], additional microsatellite sites have been identified for use in nonhuman primate studies. Microsatellite maps of the macaque MHC were recently published [Wojcechowskyj et al., 2007; Wiseman et al., 2007] in addition to a core marker set of four multiplex PCR panels comprising fifteen autosomal STR loci for genetic managements of rhesus macaque colonies [Kanthaswamy et al., 2006].

Microsatellite genotyping has already been utilized in rhesus macaque breeding groups to help define MHC haplotypes, track descent, and to differentiate chromosome configurations that would appear identical based on more limited allele-specific genotyping [Penedo *et al.*, 2005]. More recently, microsatellite analysis was instrumental in characterizing the simple genetics of Mauritian cynomolgus macaques [Wiseman *et al.*, 2007]. In order to define

MHC haplotypes in these animals, a panel of eighteen microsatellite markers, spanning the entire 5-Mb MHC region was utilized. The majority of the primers used to amplify these microsatellite markers were adapted from rhesus genomic sequences [Daza-Vamenta et al., 2004; Gourraud et al., 2004; Penedo et al., 2005]. In a cohort of more than 100 feral Mauritian cynomolgus macaques. this analysis was applied to identify the six common haplotypes that account for two-thirds of the MHC haplotypes in these animals, as described above [Wiseman et al., 2007]. Microsatellite genotyping can also be used for direct genotyping for a specific MHC class allele if a tightly linked microsatellite marker is identified. For example, the H11-9268 microsatellite marker has shown exceptional linkage to Mamu-B\*17 in a cohort of 75 Indian rhesus macaques [Wojcechowskyj et al., 2007]. In the future, this technology may assist in selective breeding of nonhuman primates by identifying homozygous animals, something that is not possible with current PCR-SSP strategies.

Identification of single nucleotide polymorphisms (SNPs) has also been aided by the completion of the rhesus macaque genome. Recently, 23,000 candidate SNPs were identified throughout the rhesus macaque genome and compiled into a web resource called *MamuSNP* that is housed at http://mamusnp.ucdavis.edu [Malhi *et al.*, 2007]. Two other studies identified and applied SNPs to population genetic analyses between Indian-origin and Chinese-origin rhesus macaques to understand the demographic history and genetic divergence of these two groups [Ferguson *et al.*, 2007; Hernandez *et al.*, 2007]. Like microsatellite analysis, genetic studies using SNPs will play an important role in establishing ancestry, directing macaque breeding programs, and helping to map and identify genes involved in complex diseases in the future.

#### **I-B-12** Concluding remarks

Nonhuman primate AIDS research is critical to HIV vaccine development and pathogenesis studies. A great deal of progress has been made in improving the nonhuman primate model in the last decade. The cellular immune responses of Indian rhesus macaques are far better defined, and additional macaque models, Burmese rhesus, pigtail, and cynomolgus, which are useful for AIDS research have been identified. However, despite this progress, our understanding of macaque immunogenetics is still is rudimentary. Much work remains to be done in identifying MHC class I and class II alleles in these macaques, in mapping immune responses, and in characterizing which responses are effective against immunodeficiency viruses. Understanding these basic issues may then facilitate pathogenesis studies, and the rational design and testing of T cell-based vaccination strategies.

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### I-B-13 Table of SIV epitopes

**Table I-B.1:** Known MHC class I epitopes and restricting molecules

Virus	Species <sup>a</sup>	Protein	Amino acid positions <sup>b</sup>	Short name	Sequence	Restricting molecule <sup>c</sup>	GenBank Acc. No. <sup>d</sup>	Reference
SIVmac239	Rhesus	Gag	149-157	LW9	LSPRTLNAW	Mamu-A*01 <sup>e</sup>	U50836	Allen et al., 2001
SIVmac239,	Rhesus	Gag	181-189	CM9	CTPYDINQM	Mamu-A*01e	U50836	Allen et al., 2001, 1998;
SIVsmE660,								Furchner et al., 1999
SIVsmH4								
SIVmac239	Rhesus	Gag	254-262	QI9	QNPIPVGNI	Mamu-A*01e	U50836	Allen et al., 2001
SIVmac239	Rhesus	Gag	372-379	LF8	LAPVPIPF	Mamu-A*01e	U50836	Allen et al., 2001
SIVmac239	Rhesus	Pol	147-156	LV10	LGPHYTPKIV	Mamu-A*01 <sup>e</sup>	U50836	Allen et al., 2001
SIVmac239	Rhesus	Pol	592-600	QV9	QVPKFHLPV	Mamu-A*01 <sup>e</sup>	U50836	Allen <i>et al.</i> , 2001
SIVmac239,	Rhesus	Pol	625-633	SV9	STPPLVRLV	Mamu-A*01 <sup>e</sup>	U50836	Allen et al., 2001; Egan et al.,
SIVmac251,								1999
SHIV-89.6,								
SHIV-HXBc2					C			
SIVmac251	Rhesus	Pol	692-700	SV9	SGPK <b>T</b> NIIV <sup>f</sup>	Mamu-A*01 <sup>e</sup>	U50836	Allen et al., 2001; O'Connor
					C			et al., 2004
SIVmac239	Rhesus	Pol	696-704	SV9	SGPK <b>A</b> NIIV <sup>f</sup>	Mamu-A*01e	U50836	Allen <i>et al.</i> , 2001
SIVmac239	Rhesus	Env	233-240	CL8	CAPPGYAL	Mamu-A*01e	U50836	Allen <i>et al.</i> , 2001
SIVmac239,	Rhesus	Env	233-241	CL9	CAPPGYALL	Mamu-A*01 <sup>e</sup>	U50836	Allen et al., 2001; Furchner
SIVsmE660,			(234-242)					et al., 1999
(SIVsmH4)		_		****		3.5	******	
SHIV-89.6,	Rhesus	Env	397-405	YI9	YAPPISGQI	Mamu-A*01 <sup>e</sup>	U50836	Egan <i>et al.</i> , 1999
SHIV-HXBc2	D.I		(20, (20	TEXT O	TUDUDUACI G	3.6 4.4018	1150026	4.11
SIVmac239	Rhesus	Env	620-628	TL9	TVPWPN <b>AS</b> L <sup>g</sup>	Mamu-A*01e	U50836	Allen <i>et al.</i> , 2001
SIVsmH4,	Rhesus	Env	626-634	TL9	TVPWPN <b>ET</b> L <sup>g</sup>	Mamu-A*01 <sup>e</sup>	U50836	Furchner et al., 1999
SIVsmE660	DI	Б	706 705	CTT10	CCDDCVEOOT	N# A #018	1150026	A11
SIVmac239	Rhesus	Env	726-735	ST10	SSPPSYFQQT	Mamu-A*01 <sup>e</sup>	U50836	Allen <i>et al.</i> , 2001; O'Connor <i>et al.</i> , 2004
SIVmac251	Rhesus	Env	729-738	ST10	SPPSYFQTHT	Mamu-A*01e	U50836	Allen et al., 2001
SIVmac239	Rhesus	Tat	28-35	SL8	<b>S</b> TPESANL <sup>h</sup>	Mamu-A*01e	U50836	Allen et al., 2001, 2000
SIVmac251	Rhesus	Tat	28-35	TL8	<b>T</b> TPESANL <sup>h</sup>	Mamu-A*01e	U50836	Allen et al., 2001
SIVmac239	Rhesus	Vif	144-152	QA9	QVPSLQYLA	Mamu-A*01e	U50836	Allen et al., 2001
SIVmac239	Rhesus	Vpx	8-18	II11	IPPGNSGEETI	Mamu-A*01e	U50836	Allen et al., 2001

Table of SIV epitopes

**Table I-B.1:** Known MHC class I epitopes and restricting molecules (cont.)

Virus	Species <sup>a</sup>	Protein	Amino acid positions <sup>b</sup>	Short name	Sequence	Restricting molecule <sup>c</sup>	GenBank Acc. No.d	Reference
SIVmac239	Rhesus	Gag	71-79	GY9	GSENLKSLY	Mamu-A*02 <sup>i</sup>	U50837	Loffredo et al., 2004; Vogel et al., 2002a
SIVmac239	Rhesus	Pol	324-332	FF9	FSIPLDEEF	Mamu-A*02 <sup>i,j</sup>	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Pol	518-526	LY9	LSQEQEGCY	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Env	296-304	RY9	RTIISLNKY	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac251	Rhesus	Env	306-313	YR8	YNLTMKCR	Mamu-A*02i	U50837	Watanabe et al., 1994
SIVmac239	Rhesus	Env	317-325	KM9	KTVLPVTIM	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Env	359-367	QY9	QTIVKHPRY	Mamu-A*02 <sup>i,j</sup>	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Env	519-528	GF10	GTSRNKRGVF	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Env	760-768	SY9	SSWPWQIEY	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Env	788-795	RY8	RTLLSRVY	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Nef	20-28	LY9	LLRARGETY	Mamu-A*02 <sup>i,j</sup>	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Nef	110-119	TM10	TMSYKLAIDM	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239,	Rhesus	Nef	159-167	YY9	YTSGPGIRY	Mamu-A*02i	U50837	Loffredo et al., 2004;
SIVmac251,								Robinson et al., 2001; Vogel
SHIV-89.6(P)								et al., 2002a
SIVmac239	Rhesus	Nef	169-177	KL9	KTFGWLWKL	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Nef	221-229	YY9	YTYEAYVRY	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Nef	248-256	LM9	LTARGLLNM	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Vif	89-97	IW9	ITWYSKNFW	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Vif	97-104	WY8	WTDVTPNY	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Vif	104-113	YY10	YADILLHSTY	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Vpr	63-71	RM9	RILQRALFM	Mamu-A*02i	U50837	Loffredo et al., 2004
SIVmac239	Rhesus	Pol	782-789	YL8	YHSNVKEL	Mamu-A*07	AF161324	Sacha et al., 2007
SHIV-HXBc2	Rhesus	Env	117-124	KP8	KPCVKLTP	Mamu-A*08	AF243179	Voss & Letvin, 1996
SIVmac239	Rhesus	Gag	178-186	SI9	SEGCTPYDI	Mamu-A*11	AF199357	Sette et al., 2005
SIVmac239	Rhesus	Pol	92-100	AL9	AERKQREAL	Mamu-A*11	AF199357	Sette <i>et al.</i> , 2005
SIVmac239	Rhesus	Pol	507-517	AI11	AEAEYEENKII	Mamu-A*11	AF199357	Sette <i>et al.</i> , 2005
SIVmac239,	Rhesus	Env	495-502	GI8	GDYKLVEI	Mamu-A*11	AF199357	Dzuris et al., 2000; Evans
(SIVppm)			(497-504)					et al., 2000, 1999; Sette et al., 2005
SIVmac239	Rhesus	Nef	124-132	KI9	KEKGGLEGI	Mamu-A*11	AF199357	Sette et al., 2005
SIVmac239	Rhesus	Vpr	13-21	RV9	REPWDEWVV	Mamu-A*11	AF199357	Sette et al., 2005

 Table I-B.1: Known MHC class I epitopes and restricting molecules (cont.)

Virus	Species <sup>a</sup>	Protein	Amino acid positions <sup>b</sup>	Short name	Sequence	Restricting molecule <sup>c</sup>	GenBank Acc. No. <sup>d</sup>	Reference
SIVppm	Rhesus	Env	575-583	KL9	KRQQELLRL	Mamu-B*03	U41825	Evans et al., 1999, 2000
SIVppm	Rhesus	Nef	136-146	AL11	ARRHRILD <b>I</b> YL <sup>k</sup>	Mamu-B*03	U41825	Dzuris <i>et al.</i> , 2000; Evans <i>et al.</i> , 2000, 1999
SIVppm	Rhesus	Nef	136-146	AL11	ARRHRILD <b>M</b> YL <sup>k</sup>	Mamu-B*03	U41825	Dzuris <i>et al.</i> , 2000; Evans <i>et al.</i> , 2000, 1999
SIVppm	Rhesus	Nef	62-69 (62-70)	QP8 (QW9)	QGQYMNTP(W)	Mamu-B*04	U41826	Dzuris <i>et al.</i> , 2000; Evans <i>et al.</i> , 2000, 1999
SIVmac239	Rhesus	Rev	12-20	KL9	KRLRLIHLL	Mamu-B*08	U41830	Loffredo et al., 2007c
SIVmac239	Rhesus	Rev	44-51	RL8	RRRWQQLL	Mamu-B*08	U41830	Loffredo et al., 2007c
SIVmac239	Rhesus	Nef	8-16	RL9	RRSRPSGDL	Mamu-B*08	U41830	Loffredo et al., 2007c
SIVmac239	Rhesus	Nef	137-146	RL10	RRHRILDIYL	Mamu-B*08	U41830	Loffredo et al., 2007c
SIVmac239	Rhesus	Nef	246-254	RL9	RRLTARGLL	Mamu-B*08	U41830	Loffredo et al., 2007c
SIVmac239	Rhesus	Vif	123-131	RL9	RRAIRGEQL	Mamu-B*08	U41830	Loffredo et al., 2007c
SIVmac239	Rhesus	Vif	172-179	RL8	RRDNRRGL	Mamu-B*08	U41830	Loffredo et al., 2007c
SHIV-HXBc2	Rhesus	Env	553-561	NA9	NNLLRAIEA	Mamu-B*12	AF243178	Voss & Letvin, 1996
SIVmac239	Rhesus	Pol	372-379	MF8	MRHVLEPF	Mamu-B*17	AF199358	Mothé et al., 2002b
SIVmac239	Rhesus	Pol	435-443	FW9	FQWMGYELW	Mamu-B*17	AF199358	Mothé et al., 2002b
SIVmac239	Rhesus	Pol	604-613	VW10	VWEQWWTDYW	Mamu-B*17	AF199358	Mothé et al., 2002b
SIVmac239	Rhesus	Env	241-251	LF11	LRCNDTNYSGF	Mamu-B*17	AF199358	Mothé et al., 2002b
SIVmac239	Rhesus	Env	816-825	LY10	LRTELTYLQY	Mamu-B*17	AF199358	Mothé et al., 2002b
SIVmac239	Rhesus	Env	830-838	FW9	FHEAVQAVW	Mamu-B*17	AF199358	Mothé et al., 2002b
SIVmac239	Rhesus	Nef	165-173	IW9	IR <b>Y</b> PKTFGW <sup>l</sup>	Mamu-B*17	AF199358	Horton <i>et al.</i> , 2001; Mothé <i>et al.</i> , 2002b
SIVppm	Rhesus	Nef	165-173	IW9	IR <b>F</b> PKTFGW <sup>l</sup>	Mamu-B*17	AF199358	Dzuris <i>et al.</i> , 2000; Evans <i>et al.</i> , 2000, 1999
SIVmac239	Rhesus	Nef	195-203	MW9	MHPAQTSQW	Mamu-B*17	AF199358	Mothé et al., 2002b
SIVmac239	Rhesus	Nef	199-207	QW9	QTSQWDDPW	Mamu-B*17	AF199358	Mothé et al., 2002b
SIVmac239	Rhesus	Vif	44-52	HW9	HFKVGWAWW	Mamu-B*17	AF199358	Mothé <i>et al.</i> , 2002b
SIVmac239	Rhesus	Vif	66-73	HW8	HLEVQGYW	Mamu-B*17	AF199358	Mothé <i>et al.</i> , 2002b
SIVmac239	Rhesus	Vif	135-143	CY9	CRFPRAHKY	Mamu-B*17	AF199358	Mothé et al., 2002b
SIVmac239	Burmese rhesus	Gag	206-216	IL11	IINEEAADWDL	90-120-Ia (hapl	lotype) <sup>m</sup>	Matano <i>et al.</i> , 2004
SIVmac239	Burmese rhesus	Gag	241-249	SW9	SSVDEQIQW	90-120-Ia (hapl	• •	Kawada <i>et al.</i> , 2006; Matano <i>et al.</i> , 2004

**Table I-B.1:** Known MHC class I epitopes and restricting molecules (cont.)

Virus	Species <sup>a</sup>	Protein	Amino acid positions <sup>b</sup>	Short name	Sequence	Restricting molecule <sup>c</sup>	GenBank Acc. No. <sup>d</sup>	Reference
SIVmac239	Burmese rhesus	Gag	373-380	AA8	APVPIPFA	90-120-Ia (hap	lotype) <sup>m</sup>	Kawada <i>et al.</i> , 2006; Matano <i>et al.</i> , 2004
SIVmac239, SIVmac251, SHIVmn229, SHIV- SF162P3	Pigtail	Gag	164-172	KP9	KKFGAEVVP	Mane-A*10 <sup>n</sup> Mane-A*16	AY557348 AY557354	Fernandez <i>et al.</i> , 2005; Smith <i>et al.</i> , 2005a,b
SIVmac239, SHIVmn229, SHIV- SF162P3	Pigtail	Gag	251-258	YP8	YRQQNPIP	Mane-A*11 Mane-A*12	AY557349 AY557350	Fernandez <i>et al.</i> , 2005; Smith <i>et al.</i> , 2005a
SIVmac239, SHIVmn229, SHIV- SF162P3	Pigtail	Gag	371-379	AF9	ALAPVPIPF	Mane-A*17	DQ886026	Fernandez et al., 2005; Loh et al., 2007; Pratt et al., 2006
SIVmac239, SHIVmn229, SHIV- SF162P3	Pigtail	Gag	28-36	KW9	KYMLKHVVW	Mane-B*10	AY557355	Fernandez <i>et al.</i> , 2005; Loh <i>et al.</i> , 2007; Smith <i>et al.</i> , 2005a
SIVmac32H- J5	Cynomologus	Gag	242-250	SM9	SVDEQIQWM	Mafa-A*02	AB154761	Geretti et al., 1997

#### Notes:

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<sup>&</sup>lt;sup>a</sup> Unless specified, rhesus designation implies rhesus macaques of Indian descent.

<sup>&</sup>lt;sup>b</sup> Positions indicated for SIVmac239 or first virus listed for each epitope unless otherwise specified. Epitopes in Pol are numbered from the open reading frame.

<sup>&</sup>lt;sup>c</sup> MHC class I molecule designations: rhesus macaque (*Macaca mulatta*; *Mamu*); cynomolgus macaque (*Macaca fascicularis*; *Mafa*); pigtail macaque (*Macaca nemestrina*; *Mane*).

<sup>&</sup>lt;sup>d</sup> The GenBank Acc. No. listed for each MHC class I restricting allele contains the most complete nucleotide sequence for that corresponding allele. Multiple entries were marked if several full length sequences were available.

<sup>&</sup>lt;sup>e</sup> The following GenBank Acc. No. are also listed for Mamu-A\*01: AJ539307 and NM 001048246.

f This CD8+ T cell epitope, with an amino acid substitution at position 5, has been identified in both SIVmac239- and SIVmac251-infected macaques.

g This CD8+ T cell epitope, with amino acid substitutions at position 7 and 8, has been identified in both SIVmac239- and SIVsmE660-infected macaques.

<sup>&</sup>lt;sup>h</sup> This CD8+ T cell epitope, with an amino acid substitution at position 1, has been identified in both SIVmac239- and SIVmac251-infected macaques.

- <sup>i</sup> The following GenBank Acc. No. is also listed for Mamu-A\*02: AJ539308.
- <sup>j</sup> Epitopes may also be restricted by Mamu-A\*01 in addition to Mamu-A\*02.
- <sup>k</sup> SIVppm is a heterogenous virus stock that contained both sequences of this Nef epitope.
- <sup>1</sup> This CD8+ T cell epitope, with an amino acid substitution at position 3, has been identified in both SIVmac239- and SIVppm-infected macaques.
- <sup>m</sup> MHC class I restriction unknown but mapped to a single MHC class I haplotype, 90-120-Ia. This haplotype consists of three *Mamu-A* alleles, (*Mamu-A120-1*, *Mamu-A120-4*, and *Mamu-A120-5*) and four *Mamu-B* alleles (*Mamu-B120-1*, *Mamu-B120-6*, *Mamu-B120-8*, and *Mamu-B120-9*).
- <sup>n</sup> The following GenBank Acc. No. is also listed for Mane-A\*10: EF010518.

**Table I-B.2:** Known MHC class II epitopes and resticting molecules

Virus	Species <sup>a</sup>	Protein	Amino acid positions <sup>b</sup>	Short name	Sequence	Restricting molecule <sup>c</sup>	GenBank Acc. No.	Reference
SIVmac239	Rhesus	Gag	102-111	QE10	QIVQRHLVVE	DRBw*606	AJ601370	Giraldo-Vela et al., 2007
			(103-112)	(IT10)	(IVQRHLVVET)			
SIVmac239	Rhesus	Gag	184-193	YV10	YDINQMLNCV	DRBw*2104	AJ601362	Giraldo-Vela et al., 2007
SIVmac239	Rhesus	Gag	197-211	GA15	QAAMQIIRDIINEEA	DRB1*0306	L27740	Giraldo-Vela et al., 2007
SIVmac239	Rhesus	Gag	260-274	GC15	GNIYRRWIQLGLQKC	DRB*w201	L27742	Dzuris et al., 2001
SHIV-89.6	Rhesus	Env	172-191	EY20	EYAFFYKLDIIPIDNDTTSY	DRB*w201	L27742	Lekutis & Letvin, 1997;
SHIV-HXBc2								Lekutis et al., 1997
SHIV-89.6	Rhesus	Env	242-261	VL20	VSTVQCTHG <b>IRPVVSTQL</b> LL <sup>d</sup>	DRB1*0406	AJ601355	Dzuris et al., 2001; Lekutis &
SHIV-HXBc2								Letvin, 1997
SHIV-89.6	Rhesus	Env	486-494	YL9	YKVVKIEPL <sup>d</sup>	DRB*w201	L27742	Dzuris et al., 2001; Lekutis &
SHIV-HXBc2								Letvin, 1997; Lekutis <i>et al.</i> , 1997
SIVmac239	Rhesus	Rev	11-23	RT13	RKRLRLIHLLHQT	DRB*w201	L27742	Dzuris et al., 2001
SIVmac239	Rhesus	Rev	13-23	RT11	RLRLIHLLHQT	DPB1*06	EF490966	Giraldo-Vela et al., 2007
SIVmac239	Rhesus	Nef	138-152	RI15	RHRILDIYLEKEEGI	DRBw*606	AJ601370	Giraldo-Vela et al., 2007
SIVmac239	Rhesus	Vpx	31-40	EL10	EINREAVNHL	DRB1*1003	AJ601356	Giraldo-Vela et al., 2007
			(32-41)	(IP10)	(INREAVNHLP)			

#### Notes:

 <sup>&</sup>lt;sup>a</sup> Unless specified, rhesus designation implies rhesus macaques of Indian descent.
 <sup>b</sup> Positions indicated for SIVmac239 or first virus listed for each epitope unless otherwise specified.
 <sup>c</sup> MHC class I molecule designation: rhesus macaque (*Macaca mulatta*; Mamu).
 <sup>d</sup> The core binding region is shown in bold [Dzuris *et al.*, 2001].

Table I-B.3: Peptide-binding motifs of macaque MHC molecules

MHC molecule	Primary anchor positions	Preferred amino acids	Tolerated amino acids	References
Mamu-A*01 <sup>a</sup>	2 <sup>b</sup> 3 <sup>b</sup> C-terminus <sup>b</sup>	S T P F L I V M	FWYNAGPLIVM TAC WYTA	Allen et al., 1998; Sidney et al., 2000
Mamu-A*02 <sup>a</sup>	2 C-terminus	T S V Y F M L W V I	L I V M A A	Loffredo <i>et al.</i> , 2004
Mamu-A*11 <sup>a</sup>	2 C-terminus	E I	D M F W L M V	Sette et al., 2005
Mamu-B*01 <sup>a</sup>	2 C-terminus	D E I F L V W	ASNGIQLTVM YM	Loffredo <i>et al.</i> , 2005
Mamu-B*03 (preliminary motif)	2 C-terminus	R L		Dzuris et al., 2000
Mamu-B*04 (preliminary motif)	2	G		Dzuris et al., 2000
Mamu-B*08 (preliminary motif)	2 C-terminus	R L I V		Loffredo <i>et al.</i> , 2007c
Mamu-B*17 <sup>a</sup>	2 C-terminus	HAMRF W	L P Q K S C W Y T G F Y	Mothé et al., 2002b
Mamu class II DR supermotif <sup>c</sup>	1 6 9	L I V M A F Y L I V M F Y S T Q A L I V M F Q		Dzuris et al., 2001

#### Notes:

<sup>&</sup>lt;sup>a</sup> A quantitative algorithm is also available for this allele at http://www.mamu.liai.org/[Peters et al., 2005a].

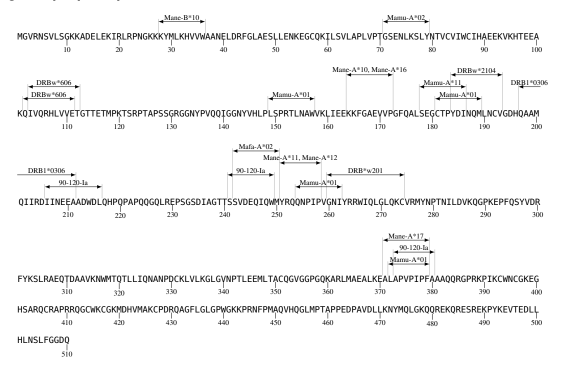
<sup>&</sup>lt;sup>b</sup> The Mamu-A\*01 peptide-binding motif requires the presence of two of the three anchor positions for binding. Peptides can bind using P2/C-terminus or P3/C-terminus anchoring spacing.

<sup>&</sup>lt;sup>c</sup> Mamu class II DR supermotif is based on data from Mamu-DRB1\*0406 and Mamu-DRB\*w201 [Dzuris *et al.*, 2001].

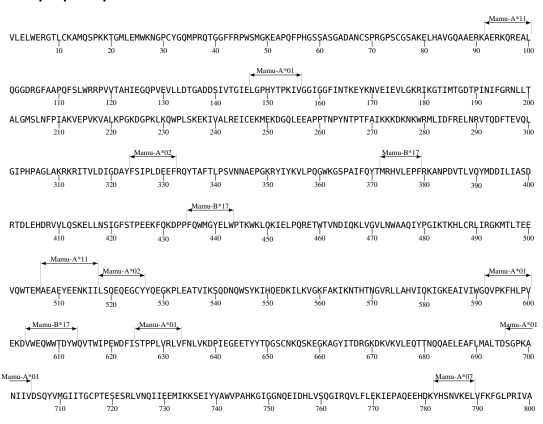
Map of SIV epitopes SIV Epitopes

#### I-B-14 Map of SIV epitopes

#### **Gag SIV Epitope Map**

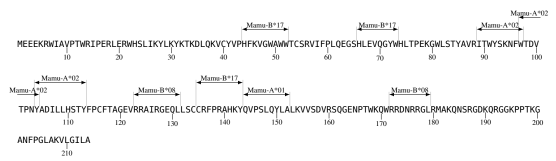


#### **Pol SIV Epitope Map**

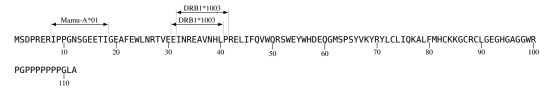




#### Vif SIV Epitope Map



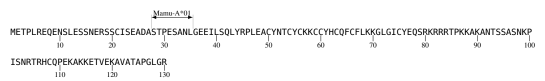
#### **Vpx SIV Epitope Map**



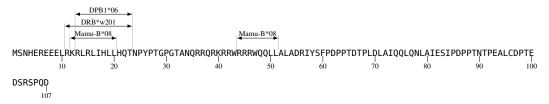
#### **Vpr SIV Epitope Map**



#### **Tat SIV Epitope Map**

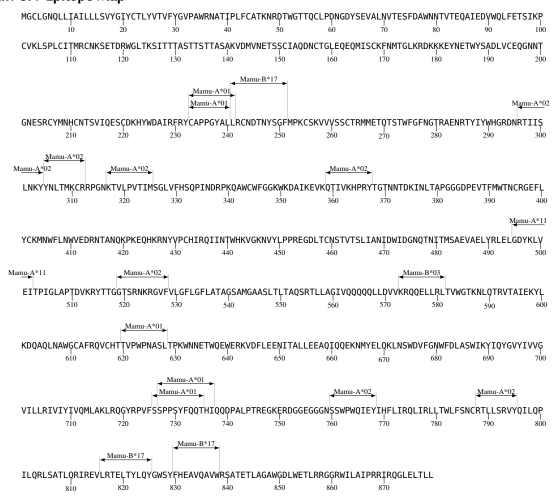


#### **Rev SIV Epitope Map**

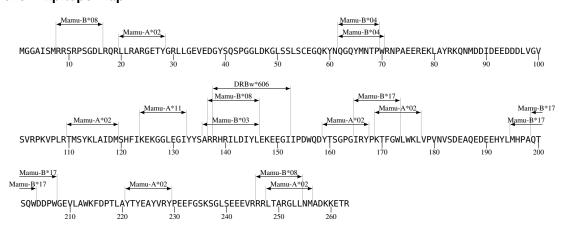


Map of SIV epitopes SIV Epitopes

#### **Env SIV Epitope Map**



#### **Nef SIV Epitope Map**



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## Part II

## **HIV CTL/CD8 + Epitopes**

### II-A

### **Summary**

This part includes tables, maps, and associated references of HIV-specific CTL epitopes arranged sequentially according to the location of the proteins in the HIV-1 genome. We attempted to make this part as comprehensive as possible, requiring that the epitope be contained within a defined region of a maximum of 30 amino acids, but not that the optimal boundaries be defined. Studies that were based on the analysis of whole proteins are described at the end of each protein section. The same epitope can have multiple entries, as each entry represents a single publication in this part of the database. For more recent updates, epitope sequence alignments, and useful search capabilities, please see our web site: http: //www.hiv.lanl.gov/content/immunology. For a concise listing of the best defined CTL epitopes, see the summary by Nicole Frahm, Caitlyn Linde and Christian Brander on page 3 in Part I of this compendium. CTL responses to proteins with no defined epitopes are listed at the end of each protein section.

Recent studies utilize multiple functions attributed to T cells to define responses, and the simple distinctions of cytotoxic T-cell and helper T-cells have become blurred as more is learned about the range of responses triggered in CD4 and CD8 positive T-cells responding to antigenic stimulus. When adding the most recent studies to the database, we have tried to place T cell responses in a reasonable manner into our traditional helper T cell and CTL parts, and to specify the assay used to measure the response in each study.

#### **II-A-1** Epitope tables

Each CTL reference has a multi-part basic entry:

HXB2 location: The viral strain HXB2 (GenBank Accession Number K03455) is used as a reference strain throughout this publication. The position of the defined epitope location relative to the sequence of the HXB2 protein is indicated. The numbering in this table corresponds to the protein maps. Because of HIV-1 variation the epitope may not actually be present in HXB2; rather, the position in HXB2 indicates the position aligned to the epitope. HXB2 was selected as the reference strain because so many studies use HXB2, and because crystal structures for HXB2-related proteins are often available. The precise positions of an epitope on the HXB2 reference strain can be readily

obtained using the interactive position locator at our web site: http://www.hiv.lanl.gov/content/hiv-db/LOCATE/locate.html.

Author location: The amino acid positions of the epitope boundaries and the reference sequence are listed as given in the primary publication. Frequently, these positions as published are imprecise, and do not truly correspond to the numbering of the sequence, but they provide a reasonable guide to the peptide's approximate location in the protein. Also, in many cases the reference sequence identification was not provided, and in such cases it is not possible to use these numbers to specify precise locations.

Epitope: The amino acid sequence of the epitope of interest as defined in the reference, based on the reference strain used in the study defining the epitope. On occasions when only the position numbers and not the actual peptide sequence were specified in the original publication, we tried to fill in the peptide sequence based on the position numbers and reference strain. If the sequences were numbered inaccurately by the primary authors, or if we made a mistake in this process, we may have misrepresented the binding site's amino acid sequence. Because of this uncertainty, epitopes that were not explicitly written in the primary publication, that we determined by looking up the reference strain and the numbered location, are followed by a question mark in the table.

**Epitope name:** If the epitope has a name attributed by the publication, it is recorded here, e.g. "SL9".

**Subtype:** The subtype under study, generally not specified for B subtype.

**Immunogen:** The antigenic stimulus of the CTL response. Often this is an HIV-1 infection. If a vaccine was used as the original antigenic stimulation, not a natural infection, this is noted separately, and additional information about the vaccine antigen is provided as available.

**Species (MHC):** The species responding and MHC or HLA specificity of the epitope.

**Donor MHC:** The HLA genotype of the individual that responded to the epitope.

**Country:** The country where the samples were obtained—generally not specified if the study was conducted in the United States.

**Assay type:** Assay used to characterize the response.

**Keywords:** Keywords are a searchable field for the web interface that is included in the T-cell sections of the

printed version to help identify entries of particular interest.

**Reference:** The primary reference (sometimes two or more directly related studies are included). Details for some of the earlier references are in Part V.

Following the entry for a given CTL epitope brief comments explain the context in which the epitope was studied and what was learned about the epitope in a given study.

#### II-A-2 HIV protein epitope maps

All HIV CTL epitopes mapped to within a region of 14 amino acids or less are indicated on the HIV protein epitope maps. The location and HLA restriction elements of CTL epitopes are indicated on protein sequences of HXB2. These maps are meant to provide the relative location of defined epitopes on a given protein, but the HXB2 sequence may not actually carry the epitope of interest, as it may vary relative to the sequence for which the epitope was defined. Epitopes with identical boundaries and HLA fields are included in the maps only once. If one laboratory determines HLA presenting molecules at the serotype level (example: A2) and another at the genotype level (example: A\*0201) both will be included in the map. MHC specificities are indicative of the host species; when no MHC presenting molecule is defined, the host species is noted.

#### **II-A-3** Alignments

To conserve space, no epitope alignments are provided in this book, but they can be generated using the CTL epitope search tool at http://www.hiv.lanl.gov/content/immunology. The master alignment files from which the epitope alignments were created are available at our web site at http://www.hiv.lanl.gov/content/hiv-db/ALIGN\_CURRENT/ALIGN-INDEX.html.

#### II-B

### **HIV CTL/CD8 + Epitope Tables**

All HIV CTL epitopes are arranged by protein position. The table entries are sorted in a nested way—first by protein, then by HXB2 start location, and finally by HLA presenting molecule. CTL reactions against proteins with undefined epitopes are listed at the end of the protein that stimulated the response.

#### II-B-1 Gag p17 CTL/CD8 + epitopes

**HXB2 Location** p17 (5–13)

**Author Location** Gag (5–13 SUMA)

Epitope ASVLSGGEL

Epitope name Gag AL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assav

Keywords dynamics, acute/early infection, characteriz-

ing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4
  T-cell counts through 7 years of followup. In contrast to more
  rapid progressors WEAU and BORI, SUMA had a broad response to 24 epitopes, with little immunodominance. Two
  peptides were somewhat more intensely recognized in acute
  infection, but this response leveled out early on.
- Only 4 epitopes acquired escape mutations in SUMA over time, and this was 1 of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

**HXB2 Location** p17 (8–18)

**Author Location** Gag (8–18)

Epitope LSGGELDRWEK

Epitope name Gag 1.2 Immunogen vaccine

infection.

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT, Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining
Keywords subtype comparisons, variant crossrecognition or cross-neutralization, memory
cells

References Amara et al. 2005

- A clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better conserved for CD8 than CD4 T cells.
- All 5 CD8 and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation. The similar reported human epitope in this case is GELDRWEKI. HLA restriction: B\*4002, B62
- Conservation across other clades: none. The form that is most common in CRF02, LSGGkLDaWEK, does not cross-react with the B clade elicited response.

HXB2 Location p17 (11-19)

**Author Location** 

Epitope GELDRWEKI

Epitope name Gag-GI9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*4002)

**Donor MHC** 01RCH46: A\*0201, A\*0217, B\*0801, B\*4002, Cw\*0303, Cw\*0701

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- This epitope was newly defined in this study.
- Patient 01RCH46 was Hispanic, on HAART, and had a viral load of 21000 and CD4 count of 623 she also recognized KETINEEAA p24(70-78), HLA B\*4002, and TAFTIPSI, RT(128-135), HLA A\*0217.
- Among HIV+ individuals who carried HLA B40, 2/5 (40%) recognized this epitope.

**HXB2 Location** p17 (11–19)

Author Location p17 (11–19)
Epitope GELDRWEKI
Immunogen HIV-1 infection
Species (MHC) human (B\*4002)
Keywords optimal epitope
References Frahm et al. 2007

HXB2 Location p17 (11–19)
Author Location p17 (11–19)
Epitope GELDRWEKI
Epitope name GI9
Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B40)

**Donor MHC** A\*30, B\*18, B\*40, Cw\*02, Cw\*05

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection, rever-

sion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs.
   Escape variants were commonly detected in maternal plasma.
   Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- The escape variant geldrwKki was detected in 10/10 maternal clones from a B40-positive mother, but was absent in all sequences from her B40-negative infant, sampled at months 2, 4, and 11, suggesting either transmission of a minority wild-type variant, or rapid reversion in the absence of continued CTL pressure.
- geldrwKki elicted lower responder cell frequencies than GEL-DRWEKI.

**HXB2 Location** p17 (11–30) **Author Location** Gag (11–30)

Epitope GELDRWEKIRLRPGGKKKYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B62)

**Donor MHC** A2, A32, B27, B62 **Assay type** Chromium-release assay

Keywords genital and mucosal immunity

References Musey et al. 2003

- CTL clones from blood, semen, cervix and rectum samples from 12 individuals were compared. CTL clones derived from blood and mucosal samples had similar high lysis efficiency, primarily perforin dependent, and TCRbeta VDJ region sequencing revealed cases of expansion of the same clone in different compartments.
- CD8+ T cell clones directed at this epitope were derived from blood and semen.

HXB2 Location p17 (12-21)

Author Location p17

Epitope ELDRWEKIRL

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human (B63)

**Assay type** CD8 T-cell Elispot - IFN $\gamma$  **Keywords** rate of progression

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate
   CTL responses early in acute HIV infection and to control HIV
   replication in the absence of antiretroviral treatment. Since
   HLA-B63 shares the epitope binding motif of HLA-B57 and
   -B58, it was shown that HLA-B63-positive individuals mounted
   CTL responses to previously identified B57-restricted epitopes,
   as well as novel, B63-restricted epitopes. Moreover, these novel
   B63-restricted epitopes can also be presented by HLA-B57 and
   -B58
- This is a putative HLA B63 epitope containing the B58 supertype binding motif embedded in a reactive peptide. There is no evidence for B57/B58 cross-presentation of this epitope.

**HXB2 Location** p17 (16–30)

**Author Location** p17 (16–30 HXB2)

**Epitope** WEKIRLRPGGKKKYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords immunodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- Responses to this peptide were detected in 19% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p17 (18–26)

**Author Location** p17 (18–26 IIIB)

Epitope KIRLRPGGK

Immunogen

Species (MHC) human (A\*0301)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an A\*0301 epitope.

**HXB2 Location** p17 (18–26)

**Author Location** 

Epitope KIRLRPGGK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301) Keywords acute/early infection References Wilson *et al.* 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

HXB2 Location p17 (18–26)
Author Location p17 (18–26 SF2)
Epitope KIRLRPGGK
Immunogen HIV-1 infection
Species (MHC) human (A\*0301)
References Altfeld *et al.* 2001a

- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- The reactive peptide p17 gag WEKIRLRPGGKKKYK contained two A\*0301-restricted epitopes, KIRLRPGGK and RL-RPGGKKK A\*0301.

HXB2 Location p17 (18–26) Author Location p17 (18–26) Epitope KIRLRPGGK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0301)

Assay type CD8 T-cell Elispot - IFNγ, CD8 T-cell Elispot granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8+ cells are found, each one constituting 30-40% of the CD8+ cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- Two of seven patients responded to this peptide with GzB producing cells, while none of the patients responded with IFN-gamma producing cells.

HXB2 Location p17 (18-26)

**Author Location** p17

Epitope KIRLRPGGK

Subtype A

Immunogen HIV-1 infection Species (MHC) human (A\*0301)

**Donor MHC** A\*0101, A\*0301, B\*0801; A\*0201, A\*3101,

B\*3501, B\*3905 Country United Kingdom.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , HLA binding

Keywords escape, acute/early infection, variant cross-

recognition or cross-neutralization

References Milicic et al. 2005

- CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. CTL escape variants were often transmitted. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient.
- A\*0301 epitopes RLRPGGKKK and KIRLRPGGK, and B\*0801 epitope GGKKKYRL, overlap. In 1 donor, the transmitted virus carried the escape form for 2 of these epitopes. The double substitution kirlrpggR results in escape from response in the donor. Similarly, the double substitution ggRkkyKl results in escape for this epitope.
- The escape mutation kirlrpggR in this epitope resulted in 74% reduction in HLA binding affinity. The other variant tested, kirlrQggR, resulted in 90% reduction.

**HXB2 Location** p17 (18-26)

Author Location p17 (18-26 IIIB)

Epitope KIRLRPGGK

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Keywords** responses in children, mother-to-infant trans-

mission, escape

References Wilson et al. 1996

- Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.
- KIRLRPGGR and RIRLRPGGR, naturally occurring variants, were found in mother and are escape mutants.

**HXB2 Location** p17 (18–26)

**Author Location** p17 (18–26)

Epitope KIRLRPGGK

Immunogen in vitro stimulation or selection

**Species (MHC)** human (A3)

**Keywords** dendritic cells

References Zarling et al. 1999

• This study compares the ability of macrophages and dendritic cells to stimulate primary responses in CD8+ lymphocytes isolated from HLA-appropriate HIV-uninfected donors using peptide-pulsed APC – the dendritic cells performed better as APC for the stimulation of primary responses.

- Strong CTL responses were elicited by the epitopes DRFYK-TLRA and GEIYKRWII when presented by either immature or mature dendritic cells – macrophages were not able to prime a CTL response against DRFYKTLRA.
- A weak response to KLTPLCVSL was stimulated using macrophages as the APC.
- No detectable response was observed for the following previously-defined HIV epitopes: KIRLRPGGK, ILKEPVHGV, IRLRPGGK, GPKVKQWPL.

HXB2 Location p17 (18–26)
Author Location Gag (18–26)
Epitope KIRLRPGGK
Immunogen HIV-1 infection
Species (MHC) human (A3)
References Brodie et al. 1999

- The ability of CTL effector cells was studied by expanding autologous HIV-1 Gag-specific CTL in vitro, and adoptive transfer.
- The transferred CTLs migrated to the lymph nodes and transiently reduced circulating productively-infected CD4+ T cells, showing that CTL move to appropriate target sites and mediate anti-viral effects.

HXB2 Location p17 (18–26)
Author Location (18–26)
Epitope KIRLRPGGK
Immunogen HIV-1 infection
Species (MHC) human (A3)
References Brodie et al. 2000

- Study tracks and quantifies in vivo migration of neo-marked CD8+ HIV-specific CTL.
- Adoptively transferred gene-marked HIV-specific CTL homed to specific lymph node sites, colocalizing within the parafollicular regions of the lymph node adjacent to cells expressing HIV tat-fusion transcripts, indicative of viral replication.
- The CTL clones expressed CCR5 and localized among HIV-1 infected cells expressing MIP-1alpha and MIP-1beta, CCchemokines produced at sites of viral replication, suggesting a possible homing mechanism.
- This study provides a methodology for tracking and studying antigen specific CTL *in vivo*.

HXB2 Location p17 (18–26)
Author Location p17 (18–26 IIIB)
Epitope KIRLRPGGK
Immunogen HIV-1 infection
Species (MHC) transgenic mouse (A3)

**Keywords** responses in children, mother-to-infant transmission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- KIRLRPGGR and RIRLRPGGR were escape mutants.
- This epitope was recognized and many escape mutants were detected in an HLA A3 transmitting mother, and was recognized but invariant in an HLA A3 non-transmitting mother.

HXB2 Location p17 (18–26) Author Location p17 (18–26 IIIB) Epitope KIRLRPGGK Immunogen HIV-1 infection Species (MHC) human (A3)

> **Keywords** review, escape **References** Goulder *et al.* 1997e; Goulder *et al.* 1997a

- HLA-identical sibling hemophiliac brothers were both infected with the same batch of factor VIII. Goulder *et al.* [1997e] is a review of immune escape that summarizes this study.
- One had a response to this epitope, the other did not. They were tested 6-8 years after infection.

HXB2 Location p17 (18–26) Author Location p17 (subtype B) Epitope KIRLRPGGK

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A3)

References Kaul et al. 2000

- 11 of 16 heavily HIV exposed but persistently seronegative sex-workers in Nairobi had HIV-specific CD8+ gamma-IFN responses in the cervix systemic CD8+ T-cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T-cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T-cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

HXB2 Location p17 (18–26) Author Location p17 (SF2) Epitope KIRLRPGGK Immunogen HIV-1 infection Species (MHC) human (A3)

**Keywords** subtype comparisons, immunodominance **References** Goulder *et al.* 2000a

- WEKIRLRPGGKKKYKLK was the target of the dominant response in Caucasoids (38%) more frequently than non-Caucasoids (12%) – 7/10 that had a dominant response to this epitope were A3, and 5/7 targeted RLRPGGKKK while 2/7 targeted KIRLRPGGK.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

**HXB2 Location** p17 (18–26) **Author Location** p17

Epitope KIRLRPGGK
Immunogen HIV-1 infection
Species (MHC) human (A3)

**Keywords** HAART, ART **References** Seth *et al.* 2001

• CTL responses were studied by tetramer staining in 41 patients with combination therapy – activated CD8+ T-cells decline as the viral load drops in response to therapy, but the overall level of antigen-specific cells capable of differentiating into effectors stays constant and new epitopes may be recognized.

HXB2 Location p17 (18–26) Author Location p17 (18–26 SF2) Epitope KIRLRPGGK Immunogen HIV-1 infection Species (MHC) human (A3)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A3+ individuals that had a CTL response to this epitope broken down by group: 3/7 group 1, 0/4 group 2, and 2/2 group 3.

HXB2 Location p17 (18–26) Author Location p17 (18–26)

Epitope KIRLRPGGK

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (A3)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- KIRLRPGGK is cross-reactive for A, B, and D clades.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** p17 (18–26)

**Author Location** p17 (JRCSF)

Epitope KIRLRPGGK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

References Severino et al. 2000

Primary HLA-A3+ CD4+ and HLA-mismatched lymphocytes from uninfected donors were infected with JRCSF after isolation then cocultured with the A3-restricted CTL clone 11504/A7 specific for KIRLRPGGK, and viral inhibition was MHC-restricted.

- Primary monocytes and monocyte-derived DC were generated from the same donors, replication of HIV-1 in these cell types was less efficient than in lymphocytes and could also be inhibited by MHC-restricted CTL.
- DC-lymphocyte cluster cultures allowed vigorous viral replication and MHC-restricted CTL viral inhibition was blunted or lost depending on the ratio of DC to CD4+ lymphocyte in the culture

HXB2 Location p17 (18-26)

Author Location p17 (18-26)

Epitope KIRLRPGGK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP).
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- All patients recognized at least 1 A3 epitope, up to 8 A3 epitopes, but none was clearly dominant.

**HXB2 Location** p17 (18–26)

Author Location p17

Epitope KIRLRPGGK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords dendritic cells

References Ostrowski et al. 2000

- The role of CD4+ T-cell help in expansion of virus-specific memory CTL was studied through co-culture ex vivo.
- Optimal expansion of HIV-1-specific memory CTL depended on CD4+ T cell help in 9 of 10 patients – CD40 ligand trimer (CD40LT) could enhance CTL in the absence of CD4+ T cell help to a variable degree in most of patients.
- Those CTL that didn't respond to CD40LT could expand with IL2 present, and IL15 produced by dendritic cells also contributes.
- The T-helper epitope used for CD4+ T-cell stimulation was the universal tetanus helper epitope TET830-843 (QYIKANSK-FIGITE)

**HXB2 Location** p17 (18–26)

Author Location p17 (18–26)

Epitope KIRLRPGGK

Epitope name A3-KK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), immunodominance, acute/early infec-

tion

References Yu et al. 2002a

- -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 8/14 HLA-A3 positive individuals had detectable A3-restricted responses during acute infection. Only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 7/8 individuals with acute responses had specific responses for this epitope.
- · KIRLRPGGK and RLRPGGKKK were the most commonly recognized HLA-A3 epitopes during acute infection, after 1 year of treatment, and after STI. RLRPGGKKK was immunodominant.

HXB2 Location p17 (18-26) Author Location p17 (18-26) Epitope KIRLRPGGK Immunogen HIV-1 infection Species (MHC) human (A3) Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4, DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay Keywords rate of progression, escape

References Geels et al. 2003

- · Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

**HXB2 Location** p17 (18–26)

**Author Location** p17 (18–26 B consensus)

Epitope KIRLRPGGK

Epitope name KK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords epitope processing, immunodominance, es-

cape, acute/early infection, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2004

• CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or • KK9 and RK9 overlap, are presented by HLA-A3, and are frequently immunodominant and involved in acute-phase primary responses. A mutation in the C-terminal flanking residue of KK9 (K to Q) (kirlrpkkg-Q) inhibits processing of the immunodominant gag KK9 epitope, resulting in rapid decline in the KK9 specific CD8+ T-cell response. At the same time it abrogates the response to RK9 through the embedded mutation rlrpggkQk. Transmission of this mutation to patients expressing HLA-A3 prevents acute-phase response to these epitopes, although the mutation can eventually revert to wild-type allowing a delayed response to the epitope.

HXB2 Location p17 (18-26)

**Author Location** p17

Epitope KIRLRPGGK

Epitope name KK9

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords review, epitope processing, escape

References Goulder & Watkins 2004

• This paper is a review of the role of CTL in HIV infection, and it uses KK9 as an example of an epitope that escapes due to a mutation beyond the epitope on the C-terminal side that probably affects proteasomal processing.

**HXB2 Location** p17 (18–26)

**Author Location** (B consensus)

**Epitope** KIRLRPGGK

Epitope name KK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A03, B14, B60, Cw3, Cw7

Country United States.

**Assay type** Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.
- One of nine individuals recognized this epitope.

**HXB2 Location** p17 (18–26)

**Author Location** p17

Epitope KIRLRPGGK

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A3)

Donor MHC A01, A03, B39, B44, Cw4, Cw6

Assay type T-cell Elispot

Keywords HIV exposed persistently seronegative

(HEPS)

References Missale et al. 2004

- HIV-specific T-cell response was tested in patients exposed to blood from a patient with highly replicating HIV; these patients were nosocomially infected with HBV, but uninfected with HIV. HIV-specific T-cell responses were directed to structural and non-structural HIV proteins in 2 patients, suggesting that the virus replicated in these patients sufficiently to prime a cell-mediated immune response that protected them from HIV infection.
- This patient responded to 3/11 HIV epitopes tested in an IFNgamma EliSpot assay. Responses were detected 16 and 20 weeks after exposure, but were lost by week 80.

HXB2 Location p17 (18–26) Author Location Gag (18–26) Epitope KIRLRPGGK Subtype B

**Immunogen** HIV-1 infection **Species** (MHC) human (A3)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** p17 (18–26) **Author Location** p24

**Epitope** KIRLRPGGK

Epitope name KK9

Immunogen

Species (MHC) (A3)

**Keywords** review, immunodominance, escape, acute/early infection, early-expressed proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

• This review discusses the importance of 3 factors that impact the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides listed as immunodominant in acute HIV-1 infection.

HXB2 Location p17 (18–26) Author Location p17 (18–26) Epitope KIRLRPGGK

Epitope name KR9 Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3)

**Donor MHC** A\*03, A\*31, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection, reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs.
   Escape variants were commonly detected in maternal plasma.
   Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- The escape variant kirlrpggR was present in 10/10 clones from an A3+ mother, was transmitted to her infant, and present in 10/10 clones at months 2 and 4, but decreased to 0/10 clones by 15 months of age in her A3- child.

HXB2 Location p17 (18-26)

Author Location p17 (18–26)

Epitope KIRLRPGGK

Immunogen HIV-1 infection

**Species (MHC)** human (A3, A\*0301, B27)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p17 (18–26)

**Author Location** p17 (18–26)

**Epitope** KIRLRPGGK

Epitope name A3-KK9 Gag

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

Country United States.

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** epitope processing, supervised treatment interruptions (STI), escape, early treatment, su-

perinfection

References Altfeld et al. 2002a

• An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response. This epitope did not vary, although the response declined over time. The authors suggest this might be due to a downstream Arg -> Thr substitution at C+2 that may impair processing.

**HXB2 Location** p17 (18–27)

Author Location (C consensus)

Epitope KIRLRPGGKK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Country South Africa.

**Assay type** CD8 T-cell Elispot - IFNγ **Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location p17 (18-27)

**Author Location** Gag

Epitope KIRLRPGGKK

Epitope name 1272

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A11)

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion, cross-presentation by different HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for KIRLRPGGKK: 36%. This
  epitope has been previously reported to be presented by A3,
  B27, B62, Bw62 and is an A11 binder, but was not confirmed
  as a CTL target in this study.

**HXB2 Location** p17 (18–27)

Author Location p17 (18-27)

**Epitope** KIRLRPGGKK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong T-helper cell responses. Only patients starting with moderately high viral load (VL) were able to reduce the VL set point. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up.
- 8/14 patients recognized this epitope.

HXB2 Location p17 (18–27) Author Location p17 (18–27 LAI) Epitope KIRLRPGGKK

Subtype B

Immunogen

Species (MHC) human (B27)

References Brander & Walker 1996

• D. Lewinsohn, pers. comm.

HXB2 Location p17 (18-27)

Author Location p17 (18-27)

Epitope KIRLRPGGKK

Immunogen HIV-1 infection

Species (MHC) human (B27)

**References** Birk *et al.* 1998b

• A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

**HXB2 Location** p17 (18–27)

**Author Location** Gag

Epitope KIRLRPGGKK

Epitope name KK10

Immunogen HIV-1 infection

Species (MHC) human (B27)

Assay type Other

Keywords rate of progression, escape

References Gao et al. 2005

- Three distinct HLA alleles known to alter the rate of AIDS progression were studied. B\*57-mediated protection occurs early in infection and the protective effect of this allele subsides after CD4 cell count drops. In contrast, B\*27 shows no protection against progression to CD4<200, but rather delays progression to an AIDS-defining illness after the CD4 counts have dropped. B\*35-mediated rapid progression to AIDS is probably a function of early decline in CD4 counts.
- KK10 escape occurs late and was shown to precede a sharp increase in viral load. The authors hypothesize the B27 benefit may arise due to enduring HLA restriction after escape from many other allotype responses has occurred.

**HXB2 Location** p17 (18-31)

Author Location p17 (18-31)

Epitope KIRLRPGGKKKYKL

Immunogen HIV-1 infection

Species (MHC) human (A3)

References Birk et al. 1998b

 A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

**HXB2 Location** p17 (18–31)

**Author Location** p17 (18–31)

Epitope KIRLRPGGKKKYKL

Immunogen HIV-1 infection

Species (MHC) human (B62)

References Lubaki et al. 1997

 82 HIV-1-specific CTL clones from 5 long-term nonprogressors were isolated and analyzed for breadth of CTL response.

- A sustained Gag, Env and Nef response was observed, and No escape mutants were observed. clones were restricted by multiple HLA epitopes, indicating a polyclonal response.
- A subject who was HLA-B62+ had CTL that recognized this peptide, and p24 LGLNKIVRMYS, and one additional unknown epitope.

HXB2 Location p17 (18-42)

Author Location p17 (18-42 IIIB)

Epitope KIRLRPGGKKKYKLKHIVWASRELE

Immunogen HIV-1 infection Species (MHC) human (A3) References Jassoy et al. 1992

· Epitope recognized by CTL clone derived from CSF.

**HXB2 Location** p17 (18–42)

**Author Location** p17 (18–42 PV22)

Epitope KIRLRPGGKKKYKLKHIVWASRELE

Immunogen HIV-1 infection Species (MHC) human (A3) References Jassoy et al. 1993

• HIV-1 specific CTLs release  $\gamma$ -IFN, and  $\alpha$ - and  $\beta$ -TNF.

**HXB2 Location** p17 (18-42)

Author Location p17 (18-42 BH10)

Epitope KIRLRPGGKKKYKLKHIVWASRELE

Immunogen HIV-1 infection Species (MHC) human (B62) References Johnson et al. 1991

• Gag CTL response was studied in three individuals.

**HXB2 Location** p17 (19–27)

**Author Location** p17 (19–27 JRCSF)

**Epitope** IRLRPGGKK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*2705)

Keywords optimal epitope References Frahm et al. 2007

• Noted by Brander to be B\*2705.

**HXB2 Location** p17 (19–27)

Author Location p17 (19–27 LAI)

Epitope IRLRPGGKK

Subtype B

Immunogen

Species (MHC) human (B27)

References Brander & Walker 1996

**HXB2 Location** p17 (19–27)

**Author Location** p17 (19–27 JRCSF)

Epitope IRLRPGGKK

Subtype B

Immunogen HIV-1 infection

Species (MHC) scid-hu mouse (B27)

Keywords escape

References McKinney et al. 1999

• Epitope-specific CTL were infused in infected human PBL-SCID mice, and transient decreases in viral load were observed, however virus was not eradicated and the HIV-specific CTL rapidly disappeared.

- Control CTL were long lived in both infected and uninfected mice, showing the rapid loss of CTL was due to target interac-

**HXB2 Location** p17 (19–27)

Author Location p17 (SF2)

Epitope IRLRPGGKK

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- · WEKIRLRPGGKKKYKLK was the target of the dominant response in Caucasoids (38%) more frequently than non-Caucasoids (12%) - 2/3 individuals that were B27+ had a dominant response to this epitope.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

**HXB2 Location** p17 (19–27)

Author Location p17 (19–27)

**Epitope** IRLRPGGKK

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Day et al. 2001

HXB2 Location p17 (19-27)

**Author Location** p17 (19–27)

Epitope IRLRPGGKK

Epitope name IK9

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords immunodominance, escape

References Goulder et al. 2001b

• This B27 epitope is generally recognized only if there is escape in the B27 dominant epitope, p24 KRWIILGLNK.

**HXB2 Location** p17 (19–27)

**Author Location** Gag

**Epitope** IRLRPGGKK

Epitope name IK9

**Immunogen** HIV-1 infection

Species (MHC) human (B27)

Donor MHC A26, B27

Assay type CD8 T-cell Elispot - IFNγ

Keywords responses in children, rate of progression, im-

munodominance, escape

References Feeney et al. 2004

• Viral load in a perinatally infected child remained low until emergence of an escape variant (kTwiilglnk) in the immunodominant CTL epitope KRWIILGLNK when the child was 7.4 years old. The emergence of this escape mutation was followed by an increase in viremia and an increase in the number of targeted CTL epitopes, measured again when the child was 9.2 years old. A low level response to IK9 was the only other epitope recognized prior to the loss of immune control and broadening of the response, and was detected in the 7.4 year sample.

HXB2 Location p17 (19-27) Author Location p17 (19-27) Epitope IRLRPGGKK Immunogen HIV-1 infection Species (MHC) human (B27)

**Donor MHC** A1, A3, B8, B35 Country United States.

Assay type Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

Keywords HAART, ART, escape, variant crossrecognition or cross-neutralization

References Casazza et al. 2005

- Evidence of continued viral evolution during HAART therapy despite low viral load was found in 1/5 patients studied.
- The dominant viral sequence was irlrpggRk, found in 12/15 clones, while the screening sequence IRLRPGGKK was found in 3/15 clones. The least frequent variant stimulated the strongest response.
- IRLRPGGKK was previously characterized as a B27 optimized epitope, which is a mismatch with patient's HLA.

**HXB2 Location** p17 (19-28)

**Author Location** Gag

Epitope IRLRPGGKKK

Epitope name 1271

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A11, A3, B62)

**Donor MHC** A03, A11, B14, B51, Cw08, Cw13

Country United States. Assay type T-cell Elispot

Keywords binding affinity, computational epitope prediction, cross-presentation by different HLA

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for IRLRPGGKKK:43% Promiscuous epitope binding to A03, B62, Bw62 and A11.

HXB2 Location p17 (20-28) Author Location p17 (20-28)

Epitope RLRPGGKKK

Immunogen HIV-1 infection Species (MHC) human (A\*03)

Keywords review, escape

References Goulder et al. 1997e: Goulder et al. 1997a

- HLA-identical sibling hemophiliac brothers were both infected with the same batch of factor VIII.
- They were tested 6-8 years after infection. One had a response to gag A3 epitope RLRPGGKKK, the other non-responder carried the sequence RLRPGGKKC.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study.

**HXB2 Location** p17 (20–28)

Author Location p17 (20–28)

**Epitope** RLRPGGKKK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes that this is an A\*0301.

HXB2 Location p17 (20-28)

**Author Location** p17

**Epitope** RLRPGGKKK

**Immunogen** HIV-1 infection

Species (MHC) human (A\*0301)

Keywords acute/early infection

References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects -3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- · No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

HXB2 Location p17 (20-28)

Author Location p17 (20–28 SF2)

Epitope RLRPGGKKK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

References Altfeld et al. 2001a

- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- The reactive peptide p17 gag WEKIRLRPGGKKKYK contained two A\*0301-restricted epitopes, KIRLRPGGK and RL-RPGGKKK A\*0301.

**HXB2 Location** p17 (20–28) **Author Location** p17 (20–28)

Epitope RLRPGGKKK

**Epitope name** RK9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0301) Donor MHC A3, A11, B35, B51

Keywords mother-to-infant transmission

References Sabbaj et al. 2002

- IFNgamma T-cell responses in breast milk of 5 HIV-1 infected women from the US and 6 from Zambia were tested using Elispot. 11/11 women responded to Gag, 8/11 responded to Pol, 7/11 women to Nef, and 2/5 women to Env peptide pools. These responses were shown to be primarily due to CD8+ Tcells in one woman, and another woman had cytolytic responses measured by Cr-release.
- Tetramer analysis of breast milk and peripheral blood samples of one volunteer showed responses to RLRPGGKKK in both compartments, 0.65% of CD3+/CD8+ cells in breast milk, and 0.22% of CD3+/CD8+ cells in peripheral blood cells.
- The frequencies of responses in the two compartments differed, and 2/4 women who responded to epitopes in Nef 101-205 and Pol 601-710 showed responses in breast milk but no detectable responses in peripheral blood cells.

HXB2 Location p17 (20-28)

**Author Location** Gag (20–28)

Epitope RLRPGGKKK

Epitope name RK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

**Donor MHC** A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Chromium-release assay

**Keywords** escape, acute/early infection, characterizing CD8+ T cells

References Bansal et al. 2005

- Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- rlrpggkkQ escape mutant showed drastically reduced avidity. The response to this peptide was not apparent until month 20, by month 32 the escape variant was present.

HXB2 Location p17 (20-28)

Author Location p17

**Epitope** RLRPGGKKK

Subtype A

**Immunogen** HIV-1 infection **Species (MHC)** human (A\*0301)

**Donor MHC** A\*0101, A\*0301, B\*0801; A\*0201, A\*3101,

B\*3501, B\*3905

Country United Kingdom.

**Assay type** CD8 T-cell Elispot - IFNγ, HLA binding **Keywords** escape, acute/early infection, variant cross-

recognition or cross-neutralization

References Milicic et al. 2005

- CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. CTL escape variants were often transmitted. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient.
- Certain escape mutations in RLRPGGKKK, rlrpggkkQ, rlrpggkkR and rlrpggkkT, resulted in nearly complete reduction in binding affinity for A\*0301. The form that was transmitted in one of the donor pairs was rlrpggRkk, and it binds to A\*0301 with comparable affinity to RLRPGGKKK. However, an escape variant was on the rise in the donor near the time of transmission, rlrpggRkT, which eventually came to fixation in the donor, illustrating the importance of the timing of transmission regarding which variant is transmitted.
- A\*0301 epitopes RLRPGGKKK and KIRLRPGGK, and B\*0801 epitope GGKKKYRL, overlap. In 1 donor, the transmitted virus carried the escape form for 2 of these epitopes. The double substitution kirlrpggR results in escape from response in the donor. Similarly, the double substitution ggRkkyKl results in escape for this epitope.

**HXB2 Location** p17 (20–28)

Author Location p17 (20-28)

Epitope RLRPGGKKK

Immunogen HIV-1 infection

Species (MHC) human (A3)

**References** Goulder *et al.* 2000c

- Two clonal CTL responses were generated in donor 021-BMC (HLA A3/3001, B42/-, Cw17/-) against different optimal versions of this epitope, one nine amino acids long, one ten.
- A previously described optimal A3 epitope overlapping this region, KIRLRPGGK, was not recognized by CTL from 021-BMC.

**HXB2 Location** p17 (20–28)

Author Location p17 (20-28)

Epitope RLRPGGKKK

Immunogen HIV-1 infection

Species (MHC) human (A3)

References Goulder et al. 1997f

 A control CTL line that reacts with this peptide was included in the study. Author Location p17 (20–28)
Author Location p17 (20–28)
Epitope RLRPGGKKK
Immunogen HIV-1 infection
Species (MHC) human (A3)
Keywords subtype comparisons

References Cao *et al.* 1997a

References Day *et al.* 2001

The consensus peptide of A, B, and D clade viruses is RLRPGThe CTL response to optimally defined CTL epitopes restricted

• The consensus peptide of C clade viruses is RLRPGGKKH and is equally reactive.

HXB2 Location p17 (20–28) Author Location p17 (SF2) Epitope RLRPGGKKK Immunogen HIV-1 infection Species (MHC) human (A3)

**Keywords** subtype comparisons, immunodominance **References** Goulder *et al.* 2000a

- WEKIRLRPGGKKKYKLK was the target of the dominant response in Caucasoids (38%) more frequently than non-Caucasoids (12%) 7/10 that had a dominant response to this epitope were A3, and 5/7 targeted RLRPGGKKK while 2/7 targeted KIRLRPGGK.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

HXB2 Location p17 (20–28)
Author Location p17 (20–28 SF2)
Epitope RLRPGGKKK
Immunogen HIV-1 infection
Species (MHC) human (A3)

**Keywords** HAART, ART, acute/early infection **References** Altfeld *et al.* 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A3+ individuals that had a CTL response to this epitope broken down by group: 5/7 group 1, 2/4 group 2, and 2/2 group 3.

HXB2 Location p17 (20–28)

Author Location p17 (20–28)

Epitope RLRPGGKKK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords rate of progression, acute/early in

**Keywords** rate of progression, acute/early infection **References** Day *et al.* 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- All patients recognized at least 1 A3 epitope, up to 8 A3 epitopes, but none was clearly dominant.

HXB2 Location p17 (20–28) Author Location p17 (20–28) Epitope RLRPGGKKK Epitope name RK9 Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords acute/early infection

References Goulder et al. 2001b

- Responses to this dominant A3-restricted Gag epitope are present during the time of decreasing viral load in acute infection.
- Mutations in this epitope were observed in autologous clones of subjects who were A3-positive with a higher frequency than those who were A3-negative (P=0.0002)
- These mutations are being sexually transmitted in adult infections

HXB2 Location p17 (20–28)

**Author Location** 

Epitope RLRPGGKKK

**Epitope name** Gag-RK9

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (A3)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA A03, 7/20 (35%) recognized this epitope.

**HXB2 Location** p17 (20–28)

Author Location p17 (20–28)

Epitope RLRPGGKKK

Epitope name A3-RK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), immunodominance, acute/early infec-

tion

References Yu et al. 2002a

- -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06 was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 8/14 HLA-A3 positive individuals had detectable A3-restricted responses during acute infection. Only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 7/8 individuals with acute responses had specific responses for this epitope.
- · KIRLRPGGK and RLRPGGKKK were the most commonly recognized HLA-A3 epitopes during acute infection, after 1 year of treatment, and after STI. RLRPGGKKK was immunodominant during acute infection and throughout the study period in the 5/6 individuals who targeted it.

HXB2 Location p17 (20-28) **Author Location** Gag (LAI)

Epitope RLRPGGKKK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3)

Keywords class I down-regulation by Nef

References Lewinsohn et al. 2002

- CTL kill targets through releasing perforin, that forms pores in the plasma membrane, and granzymes, that induce apoptosis.
- Vpr is capable of arresting infected cells in the G2 phase, and it was hypothesized that Vpr may inhibit CTL-mediated apoptosis because it interacts with the granzyme B molecular complex.
- Vpr expression in the target cell did not inhibit epitope specific lysis - neither perforin or granzyme mediated events were inhibited, as measured by a Chromium release assay and a TUNEL assay.
- In contrast, deletion of Nef, which is thought to protect primary HIV infected cells by down-regulating cell-surface expression of MHC class I complexes, increased the susceptibility of HIV-1 infected cells to CTL mediated killing 2-fold using the TUNEL assay.

**HXB2 Location** p17 (20–28)

Author Location p17

**Epitope** RLRPGGKKK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A3, A11, B35, B51

Keywords mother-to-infant transmission

References Sabbaj et al. 2002

• IFNgamma T-cell responses in breast milk of 5 HIV-1 infected women from the US and 6 from Zambia were tested with using Elispot. 11/11 women responded to Gag, 8/11 responded to Pol, 7/11 women to Nef, and 2/5 women to Env peptide pools. These responses were shown to be primarily due to CD8+ T-cells in one woman, and another woman had cytolytic responses measured by Cr-release.

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or T-cells in breast milk from a volunteer who was HLA A3, A11, B35, B51 induced IFNgamma after stimulation with a peptide that carries known A3 epitope RLRPGGKKK.
  - The frequencies of responses in the two compartments differed, and 2/4 women that responded to epitopes in Nef 101-205 and Pol 601-710 showed responses in breast milk but no detectable responses in peripheral blood cells.

HXB2 Location p17 (20-28)

Author Location p17 (20-28)

**Epitope** RLRPGGKKK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A\*0201, A3, B44, B57, Cw5, Cw6; A1, A3, B7, B14, Cw\*0702, Cw\*0802; A1, A3, B8,

B35; A1, A3, B8, B62, Cw3, Cw7

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords acute/early infection, early-expressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- This epitope was recognized in four individuals during early infection, each time presented by A3.
- All HIV-1 proteins except Vpu were recognized, and responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

HXB2 Location p17 (20-28)

Author Location p17 (20–28)

**Epitope** RLRPGGKKK

Epitope name A3-RK9 Ga9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Assay type CD8 T-cell Elispot - IFNγ

Keywords supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

· An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.

• The second infecting strain had the variant rlrpggkkT. The CTL response declined over time, and the response to the second variant was lower than to the first one throughout.

HXB2 Location p17 (20-28) Author Location p17 (20–28) Epitope RLRPGGKKK Immunogen HIV-1 infection Species (MHC) human (A3) DR8, DQ7

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 5 epitopes from this individual that varied over time where the internal mutations became fixed; 4/5 epitopes (all except p17 RLRPGGKKK, this eptiope) showed a dramatic decrease in CTL activity against the wild type epitope as the mutation arose. The rlrpggkkR variant was found at 47 and 120 months post-seroconversion.

HXB2 Location p17 (20-28)

**Author Location** Gag

Epitope RLRPGGKKK Immunogen HIV-1 infection

Species (MHC) human (A3)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords HIV exposed persistently seronegative (HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- 0/5 HLA A3+ infection-resistant men, and 0/3 preseroconversion men who went on to become infected, reacted to this epitope.

**HXB2 Location** p17 (20–28)

**Author Location** Gag (20–28)

Epitope RPRPGGKKK Immunogen HIV-1 infection

Species (MHC) human (A3)

Assay type Cytokine production, proliferation, CD8 Tcell Elispot - IFNy, Tetramer binding, Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

Keywords HAART, ART, memory cells, characterizing

CD8+ T cells

References Daniel et al. 2004

• CD4+ and CD8+ responses in chronically HIV-1 infected patients on HAART were weak with decreased polyclonality. Only 33% of patients had CD4+ T-cells that could proliferate, and only 22% had HIV-specific CD8+ T-cell responses, and those rare responses showed low perforin levels and persistent expression of CD27, indicating incomplete differentiation and loss of lytic function.

HXB2 Location p17 (20-28)

Author Location p17 (20–28 B consensus)

Epitope RLRPGGKKK

Epitope name RK9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

Keywords immunodominance, escape, acute/early infection, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2004

• KK9 and RK9 overlap, are presented by HLA-A3, and are frequently immunodominant and involved in acute-phase primary responses. A mutation in the C-terminal flanking residue of KK9 (K to Q) (kirlrpkkg-Q) inhibits processing of the immunodominant gag KK9 epitope, resulting in rapid decline in the KK9 specific CD8+ T-cell response. At the same time it abrogates the response to RK9 through the embedded mutation rlrpggkQk. Transmission of this mutation to patients expressing HLA-A3 prevents acute-phase response to these epitopes, although the mutation can eventually revert to wild-type allowing a delayed response to the epitope.

HXB2 Location p17 (20-28)

**Author Location** (B consensus)

**Epitope** RLRPGGKKK

Epitope name RK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A02, A03, B08, B62, Cw7, Cw10; A01, A03,

B08, B14, Cw7, Cw8

Country United States.

perforin expression was found.

Assav type Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory

cells, characterizing CD8+ T cells References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
- 2/9 individuals recognized this epitope, presented by HLA-A3.

HXB2 Location p17 (20-28)

**Author Location** Gag

**Epitope** RLRPGGKKK

Epitope name RK9

Subtype B

Immunogen HIV-1 infection **Species (MHC)** human (A3)

Assav type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 9, rrlpggkkR, was found in the most polymorphic residue in the epitope. This was shared between clades B and C.

HXB2 Location p17 (20-28)

Author Location Gag (20-28 BRU)

Epitope RLRPGGKKK

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (A3)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 0/9 CRF02\_AG-infected Ivorians, and 2/9 B-infected French subjects.

HXB2 Location p17 (20-28)

**Author Location** p24

**Epitope** RLRPGGKKK

Epitope name RK9

**Immunogen** 

Species (MHC) (A3)

Keywords review, immunodominance, escape, acute/early infection, early-expressed

proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

• This review discusses the importance of 3 factors that impact the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides listed as immunodominant in acute HIV-1 infection.

**HXB2 Location** p17 (20–28)

**Author Location** Gag

Epitope RLRPGGKKK

Epitope name 1332

Subtype multiple

Immunogen HIV-1 infection

**Species (MHC)** human (A3, A\*0301, B62, B42)

Donor MHC A03, A23, B49, B57; A03, A24, B27, B57,

Cw13, Cw18; A03, A26, B08, B52

Country United States.

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope predic-

tion, immunodominance, cross-presentation

by different HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for RLRPGGKKK: 34% Promiscuous epitope binding to A03, A0301, B62, Bw62, B42. Immunodominant epitope.

**HXB2 Location** p17 (20–28)

Author Location p17 (20-28)

**Epitope** RLRPGGKKK

Immunogen HIV-1 infection

Species (MHC) human (A3, A30, B42, B62)

**Donor MHC** A2, A31, B51, B58w4

Country United States.

Assay type Intracellular cytokine staining, Flow cytomet-

ric T-cell cytokine assay

Keywords HAART, ART, escape, variant cross-

recognition or cross-neutralization

References Casazza et al. 2005

- Evidence of continued viral evolution during HAART therapy despite low viral load was found in 1/5 patients studied.
- The majority of viral sequences prior to therapy were rlrpggkkQ. At week 14 of therapy a major change in the viral quasispecies occurred: the variants present were found to be rlrpggkkK (14/16 clones) and rlrpggkkR (2/16 clones), both well recognized by HIV-specific CD8 T cells. At week 19, the quasispecies reverted back to the less well-recognized rlrpggkkQ variant.

**HXB2 Location** p17 (20–28)

**Author Location** p17 (20–28)

**Epitope** RLRPGGKKK

Immunogen HIV-1 infection

Species (MHC) human

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- Three of the four individuals that responded to SLYNTVATL recognized HIV epitopes, and one individual who was A\*0201, A31 and B51 and B58w4 recognized this epitope (previously described as HLA A3.1), as well as one other.

HXB2 Location p17 (20–29)
Author Location p17 (20–29)
Epitope RLRPGGKKKY
Immunogen HIV-1 infection
Species (MHC) human (A\*0301)
References Brander & Walker 1995

• Unpublished, C. Jassoy and Beatrice Culman, pers. comm.

HXB2 Location p17 (20–29)
Author Location p17 (20–29 LAI)
Epitope RLRPGGKKKY
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (A\*0301)

**References** Wilkens & Ruhl 1999 • Pers. comm., B. Wilkens and D. Ruhl.

HXB2 Location p17 (20–29) Author Location p17 (20–29 LAI) Epitope RLRPGGKKKY Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0301) Keywords optimal epitope

References Frahm *et al.* 2007
• C. Brander notes this is an A\*0301 epitope.

HXB2 Location p17 (20–29)
Author Location (C consensus)
Epitope RLRPGGKKHY
Subtype C
Immunogen HIV-1 infection

Species (MHC) human (A\*3002)
Country South Africa.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- RLRPGGKKHY is an optimal epitope.

HXB2 Location p17 (20–29)
Author Location p17 (20–29)
Epitope RLRPGGKKKY
Immunogen HIV-1 infection
Species (MHC) human (A3)
References Goulder et al. 2000c

- Two clonal CTL responses were generated in donor 021-BMC (HLA A3/3001, B42/-, Cw17/-) against different optimal versions of this epitope, one nine amino acids long, one ten.
- A previously described optimal A3 epitope overlapping this region, KIRLRPGGK, was not recognized by CTL from 021-BMC.

HXB2 Location p17 (20–29) Author Location p17 (20–29) Epitope RLRPGGKKKY Immunogen HIV-1 infection

Species (MHC) human (A3, A30, B42, B62)

**Donor MHC** A2, A3, B7, B44 **Country** United States.

Assay type Intracellular cytokine staining, Flow cytomet-

ric T-cell cytokine assay

Keywords HAART, ART, escape, variant cross-

recognition or cross-neutralization

References Casazza et al. 2005

- Evidence of continued viral evolution during HAART therapy despite low viral load was found in 1/5 patients studied.
- The epitope RLRPGGKKKY was invariant (18/18 sequences) prior to therapy in the patient that recognized it.

HXB2 Location p17 (20–29) Author Location p17 (20–29) Epitope RLRPGGKKKY Epitope name RY10

**Immunogen** HIV-1 infection **Species** (MHC) human (A30)

**Donor MHC** A\*24, A\*30, B\*39, B\*47, Cw\*12, Cw\*17; A\*30, B\*18, B\*40, Cw\*02, Cw\*05

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection, reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs.
   Escape variants were commonly detected in maternal plasma.
   Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- RLRPGGKKKY was recognized in 2 mothers, and is an A\*30 epitope. The variant RLRPGGKKqY was found in 9/10 of 1 mother's sequences. This form was transmitted to her child, and 10/10 clones were this variant at months 2 and 6 in the infant; by month 12, 9/10 were RLRPGGKKqY. RLRPGGKKrY was the form found in the other mother. The variant gradually diminished in frequency in her child, 10/10 sequences at 2 months, 9/10 at 4 months, and 6/10 at 12 months.

HXB2 Location p17 (20–29)
Author Location p17 (20–29)
Epitope RLRPGGKKKY
Immunogen HIV-1 infection
Species (MHC) human (A30, A\*0301)
Keywords immunodominance
References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INFγ responses to other epitopes.
- 1/11 of the A2+ individuals was A30, and one was A3, and both responded to RLRPGGKKKY.

The A2+ A3 individual also reacted with two other A3.1 epitopes.

HXB2 Location p17 (20–29)
Author Location p17 (20–29 IIIB)
Epitope RLRPGGKKKY
Immunogen HIV-1 infection
Species (MHC) human (B42)

Keywords responses in children, mother-to-infant trans-

mission

References Wilson et al. 1996

- Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.
- RLRPGGKKRY, a naturally occurring variant, was found in non-transmitting mother and is recognized.
- Binds HLA-A3 and Bw62 as well.

HXB2 Location p17 (20–29) Author Location p17 (20–29) Epitope RLRPGGKKKY Immunogen HIV-1 infection Species (MHC) human (B42, B62) References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location p17 (20–29) Author Location p17 (20–29 LAI) Epitope RLRPGGKKKY Subtype B

Subtype 1 Immunogen

Species (MHC) human (B62)

Keywords review

References McMichael & Walker 1994

- · Review of HIV CTL epitopes.
- Also P. Johnson, pers. comm.

HXB2 Location p17 (20–29)
Author Location p17 (20–29)
Epitope RLRPGGKKKY
Immunogen HIV-1 infection
Species (MHC) human (B62)
References Brodie et al. 2000

- Study tracks and quantifies *in vivo* migration of neo-marked CD8+ HIV-specific CTL.
- Adoptively transferred gene-marked HIV-specific CTL homed to specific lymph node sites, colocalizing within the parafollicular regions of the lymph node adjacent to cells expressing HIV tat-fusion transcripts, indicative of viral replication.
- The CTL clones expressed CCR5 and localized among HIV-1 infected cells expressing MIP-1 $\alpha$  and MIP-1 $\beta$ , CC-chemokines produced at sites of viral replication, suggesting a possible homing mechanism.
- This study provides a methodology for tracking and studying antigen specific CTL in vivo.

HXB2 Location p17 (20–29) Author Location Gag (20–29) Epitope RLRPGGKKKY Epitope name RY10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B62)

**Donor MHC** A\*01, A\*11, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** escape, variant cross-recognition or cross-neutralization

**References** Koibuchi *et al.* 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- The RY10 variant RLRPGGrKKY was the only form of the epitope detected over a 5 year time period in this person. Elispot reactions were stronger to the autologous form than to RLRPGGKKKY, the B clade consensus form.

**HXB2 Location** p17 (20–30)

Author Location p17 (SF2)

Epitope RLRPGGKKKYK

Immunogen HIV-1 infection

Species (MHC) human

 $\label{lem:keywords} \textbf{Keywords} \ \ \text{subtype comparisons, immunodominance}$ 

References Goulder et al. 2000a

- WEKIRLRPGGKKKYKLK was the target of the dominant response in Caucasoids (38%) more frequently than non-Caucasoids (12%) – the dominant response in a Haitian immigrant living in Boston who was HLA A24/29 B7/B44 Cw6/7 was to this epitope, although the restricting element was not determined.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

**HXB2 Location** p17 (20–35)

Author Location p17 (90–105 SF2)

Epitope CLRPGGKKKYKLKHIV

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.

• The responding subject was HLA A-2, A-24, B-13, B-35.

**HXB2 Location** p17 (21–35)

**Author Location** Gag

Epitope LRPGGKKKYKLKHIV

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords TCR usage

References Weekes et al. 1999b

- Peptide 703.3: Almost all CD8+ T cells are CD28+ at birth, and the proportion of CD28-CD8+ cells increases with age this study examines the contribution of CD8+CD28- cells to CTL memory pools for CTL clones specific for two persistent human viruses, CMV and HIV clones were found to be similarly distributed in the CD28 depleted cell population.
- HIV CTL responses to 3 Env and 2 Gag peptides were studied.
- The clonal composition of the TCR  $V\beta$  responses was studied and was found to be highly focused, with one TCR beta-chain sequence tending to dominate the peptide-specific response clones to this epitope were  $V\beta$ 13.1 and  $V\beta$ 5.2.

**HXB2 Location** p17 (21–35)

Author Location p17 (21-35)

Epitope LRPGGKKKYKLKHIV

Immunogen

Species (MHC) human (B8)

References Nixon & McMichael 1991

• Two CTL epitopes defined (see also p24(191-205))

**HXB2 Location** p17 (21–35)

**Author Location** p17 (21–35)

Epitope LRPGGKKKYKLKHIV

Immunogen HIV-1 infection

Species (MHC) human (not B8) References van Baalen *et al.* 1996

• Unknown HLA specificity, but not B8.

**HXB2 Location** p17 (21–35)

**Author Location** Gag

Epitope LRPGGKKKYKLKHIV

Immunogen HIV-1 infection

Species (MHC) human

References Weekes et al. 1999a

 Peptide 703.3: Memory CTL specific for HIV-1 may contribute to oligoclonal expansions within the CD57+ CD28- CD8+ CTL populations.

**HXB2 Location** p17 (21–35)

**Author Location** p17 (91–105 SF2)

Epitope LRPGGKKKYKLKHIV

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A1, A2, B50, B57.

**HXB2 Location** p17 (21–35)

**Author Location** p17 (24–31)

Epitope LRPGGKKKYRLKHLV

Subtype A, D

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*6601, A\*6801, B\*5301, B\*5802; A\*3002,

A\*6801, B\*5703, B\*5802

Country Uganda.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, variant cross-

recognition or cross-neutralization

References Barugahare et al. 2005

- T-cell immune responses were examined in HIV-1 infected Ugandans. Similar levels of cross-clade responses for Gag, Env and Nef were observed. Higher frequencies of responses in conserved regions were found for Gag within a single clade, while areas with higher viral sequence variation had fewer detectable responses. Differential interclade sequence homologies between the Gag regions did not affect the respective level of cross-clade recognition.
- The sequence contains a known B8 epitope, but the subjects recognizing it were B8-negative. The autologous viral sequence was lrpggkkkyKlkhlv, and the peptide was recognized.

**HXB2 Location** p17 (21–40)

**Author Location** p17 (21–40 subtype A)

Epitope LRPGGKKKYRLKHLVWASRE

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (Cw4)

**Keywords** subtype comparisons

**References** Dorrell *et al.* 1999

- CTL responses in three individuals with non-clade B infections were studied, 2 with subtype A infections, 1 with subtype C – their infections all originated in East Africa.
- This epitope was defined in an A subtype infection the B clade variant (LRPGGKKKYKLKHIVWASRE) has two mutations relative to the A subtype form, and the CTLs from this patient were not A-B cross-reactive.

**HXB2 Location** p17 (22–30)

Author Location p17 (22–30)

Epitope RPGGKKRYM

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*35, Cw\*0602)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope

prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of

Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

• This is a novel epitope that may be subtype C-specific.

HXB2 Location p17 (22–31) Author Location Gag (22–31) Epitope RPGGKKRYKL Immunogen HIV-1 infection Species (MHC) human (B7) References Jin et al. 2000b

- This B7 epitope is one of three subdominant CTL responses detected in a long-term non-progressor.
- A dominant B7 epitope was defined using conventional methods, and three additional sub-dominant HLA B7 epitopes were defined by first using a non-anchor based strategy, EpiMatrix, to identify 2078 possible epitopes in the autologous HIV-1, followed by B7 anchor residue prediction to narrow the set to 55 peptides for experimental testing.

 $\textbf{HXB2 Location} \ \ p17 \ (22\text{--}31)$ 

Author Location p17

Epitope RPGGKKKYKL

Subtype B, D

Immunogen HIV-1 infection

Species (MHC) human (Cw4)

Donor MHC A23, A34, B44, B53, Cw4, Cw6

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence had an K5N change, RPGGnKKYKL.

**HXB2 Location** p17 (24–31)

Author Location p17

Epitope GGKKKYKL

Epitope name GL8

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

**Donor MHC** A\*0101, A\*0301, B\*0801; A\*0201, A\*3101,

B\*3501, B\*3905

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFNγ, HLA binding

**Keywords** escape, acute/early infection, variant cross-recognition or cross-neutralization

References Milicic et al. 2005

- Escape mutation ggRkkyKl in this epitope, GGKKKYRL, resulted in failure of recognition by CTLs, and the ggkkQyRl mutations resulted in 82% reduction in HLA binding affinity.
- A\*0301 epitopes RLRPGGKKK and KIRLRPGGK, and B\*0801 epitope GGKKKYRL, overlap. In 1 donor, the transmitted virus carried the escape form for 2 of these epitopes. The double substitution kirlrpggR results in escape from response in the donor. Similarly, the double substitution ggRkkyKl results in escape for this epitope.

**HXB2 Location** p17 (24–31)

Author Location p17 (24-31)

Epitope GGKKKYKL

Immunogen

Species (MHC) human (B8)

References Goulder et al. 1997g

- The crystal structure of this peptide bound to HLA-B8 was used to predict new epitopes and the consequences of epitope variation
- The predictions were experimentally confirmed.
- The anchors for HLA-B8 epitopes, as defined by peptide elution data, are P3 (K), P5 (K/R), and P8 (L).
- Structural data suggests that a positive charge at P5 is essential, but that the constraints on P3 may be less severe.
- Small hydrophobic residues at P2 may be favorable for binding.
- A spacious F-pocket favors mid-sized hydrophobic residues in the C-term anchor.

**HXB2 Location** p17 (24–31)

Author Location p17 (24–31 SF2)

Epitope GGKKKYKL

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Keywords** subtype comparisons

References McAdam et al. 1998

CTL from a patient infected with clade B virus did not recognize Ugandan variants of this epitope.

**HXB2 Location** p17 (24–31)

Author Location p17 (24-31 LAI)

Epitope GGKKKYKL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords TCR usage

References Reid et al. 1996

- The variants 7R: GGKKKYRL, 7Q: GGKKKYQL, 5R: GGKKRYKL, and 3R: GGRKKYKL, were studied.
- Crystal structures were obtained to study these peptides in the context of HLA-B8, and CTL binding and activity were determined.
- 3R has been detected in 3 patients, and it abolishes recognition causing extensive conformational changes upon binding including MHC main chain movement.
- 7Q and 7R alter the TCR exposed surface, and retain some recognition.
- Reactivity of 5R depends on the T cell clone, this amino acid is embedded in the C pocket of B8 when the peptide is bound.

• Optimal peptide is 8-mer, not 9-mer, and positions 3, 5, and 8 are the anchor residues.

**HXB2 Location** p17 (24–31) **Author Location** p17 (24–31 LAI)

Epitope GGKKKYKL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8) References Price *et al.* 1997

• A weak CTL response to the index peptide was observed in an HLA-B8+ infected individual.

 Sequences from the earliest available time point showed that a variant at position 5, an anchor residue, GGKKQYKL, was present.

**HXB2 Location** p17 (24–31)

**Author Location** p17

Epitope GGKKKYKKL
Immunogen HIV-1 infection
Species (MHC) human (B8)
Keywords HAART, ART
References Seth et al. 2001

 CTL responses were studied by tetramer staining in 41 patients with combination therapy – activated CD8+ T-cells decline as the viral load drops in response to therapy, but the overall level of antigen-specific cells capable of differentiating into effectors stays constant and new epitopes may be recognized.

HXB2 Location p17 (24–31) Author Location p17 (24–31 SF2) Epitope GGKKKYKL

Immunogen HIV-1 infection
Species (MHC) human (B8)

**Keywords** HAART, ART, acute/early infection **References** Altfeld *et al.* 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B8+ individuals that had a CTL response to this epitope broken down by group: 0/3 group 1, 1/3 group 2, and 2/2 group 3.

**HXB2 Location** p17 (24–31)

**Author Location** p17 (24–31)

Epitope GGKKKYRL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B8)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** p17 (24–31)

Author Location p17 (24-31)

Epitope GGKKKYKL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Day et al. 2001

 B8-restricted CTL accounted for about 1/3 of the total CTL response in one individual.

**HXB2 Location** p17 (24–31)

**Author Location** p17

Epitope GGKKKYKL
Immunogen HIV-1 infection
Species (MHC) human (B8)

 $\textbf{Keywords} \ \ \text{binding affinity, review, subtype comparisons,}$ 

epitope processing, escape

References McMichael & Hanke 2002

 CTL response-eliciting vaccines are reviewed. The natural epitope interactions with the HLA class I presenting molecules and T-cell receptors are described, using the structure of this epitope, taken from Reid et al. [1996], as an example.

**HXB2 Location** p17 (24–31)

Author Location (B consensus)

Epitope GGKKKYKL

Epitope name GL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A01, A03, B08, B14, Cw7, Cw8

Country United States.

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** p17 (24–31)

**Author Location** p17

Epitope GGKKKYKL

Subtype B, D

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A1, A1, B8, B55, Cw3, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence had an L8M change, GGKKKYKm.

**HXB2 Location** p17 (24–31)

Author Location Gag (24-31 BRU)

Epitope GGKKKYKL

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (B8)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 1/9 CRF02\_AG-infected Ivorians, and 0/9 B-infected French subjects.

**HXB2 Location** p17 (24–31)

Author Location p17 (24-31)

Epitope GGKKKYKL

Epitope name GL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A\*03, A\*31, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant trans-

mission, escape, characterizing CD8+ T cells, reversion, viral fitness

reversion, vital littless

References Sanchez-Merino et al. 2005

CD8 T-cell responses were examined in mother-infant pairs.
 Escape variants were commonly detected in maternal plasma.
 Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting

- that infants are capable of generating virus-specific CD8 T-cell responses.
- Variant sequence ggRkkykl was present in 10/10 clones from a B8-positive mother, but decreased to 0/10 clones by 15 months of age in her B8-negative child.
- The variant ggRkkykl was present in 10/10 clones from a B8+ mother, was transmitted to her infant, and present in 10/10 clones at months 2 and 4, but decreased to 0/10 clones by 15 months of age in her B8- child.

**HXB2 Location** p17 (24–31)

**Author Location** p17 (24–31 HXB2)

Epitope GGKKKYKL

Epitope name GL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A\*0101, A\*0201, B\*0801, B\*50, Cw\*0602,

Cw\*0701

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, reversion, viral fit-

ness, optimal epitope, HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- GGKKKYKf presumed escape variant was transmitted to an HLA-B8 negative reipient from an HLA-B8 positive donor. Reversions were found later, with loss of CTL pressure and hypothesised gain of replicative fitness.

**HXB2 Location** p17 (24–32)

Author Location p17 (24-32 LAI)

Epitope GGKKKYKLK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

**Keywords** optimal epitope

**References** Frahm *et al.* 2007

• C. Brander notes epitope to be presented by B\*0801.

**HXB2 Location** p17 (24–32)

**Author Location** p17 (24–32 LAI)

Epitope GGKKKYKLK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**References** Sutton *et al.* 1993

• Exploration of HLA-B8 binding motif through peptide elution.

**HXB2 Location** p17 (24–32)

**Author Location** p17 (24–32 LAI)

Epitope GGKKKYKLK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8)

Keywords epitope processing

References Rowland-Jones et al. 1993

Study of an individual with partially defective antigen processing.

HXB2 Location p17 (24–32)
Author Location p17 (24–32)
Epitope GGKKKYKLK
Immunogen HIV-1 infection
Species (MHC) human (B8)

References Klenerman et al. 1994

 Naturally occurring variants GGKKKYQLK and GGKKRYRLK may act as antagonists.

HXB2 Location p17 (24–32) Author Location p17 (24–32) Epitope GGKKKYKLK Immunogen HIV-1 infection Species (MHC) human (B8)

References Klenerman et al. 1995

 Naturally occurring antagonist GGKKKYQLK found in viral PBMC DNA and RNA.

HXB2 Location p17 (24–32) Author Location p17 (24–32) Epitope GGKKKYKLK Immunogen HIV-1 infection Species (MHC) human (B8) Keywords escape

References Nowak et al. 1995

• Longitudinal study of CTL response and immune escape – the variant GGRKKYKLK binds to HLA-B8 but is not reactive.

HXB2 Location p17 (24–32) Author Location p17 (24–32) Epitope GGKKKYKLK Immunogen HIV-1 infection Species (MHC) human (B8) References Dyer et al. 1999

- CTL specific responses were measured over a 1.3 to 1.5 year period in members of the Sydney Blood Bank Cohort (SBBC) who had been infected with a natural attenuated strain of HIV-1 that was Nef-defective.
- Some of these patients had prolonged high levels of CTL effector and memory cells despite low viral load.

**HXB2 Location** p17 (24–32)

Author Location p17

Epitope GGKKKYKLK

Immunogen

Species (MHC) human (B8)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive, and the B35 allele seems to be protective.

 HIV-2 sequence: GGKKKYKMK – no cross-reactivity Phillips et al. [1991].

**HXB2 Location** p17 (24–32) **Author Location** p17 (24–32)

Epitope GGKKKYKLK

Epitope name GGK

Immunogen HIV-1 infection Species (MHC) human (B8)

**Keywords** HAART, ART, supervised treatment interruptions (STI), immunodominance, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
  - This epitope was recognized by 1/7 study subjects that were HLA-B8+.
- Patient SC12(HLA A1, B8/39, Cw0701/0702, DR2/3, DR51/52, DQ2/6) had sustained therapy started during acute infection and maintained an immunodominant response to FLKEKGGL throughout and minor responses to GEIYKR-WII, DCKTILKAL, GGKKKYKLK – GEIYKRWII and GGKKKYKLK responses were stimulated by a brief period off therapy.

HXB2 Location p17 (24-32)

Author Location p17

Epitope GGKKKYKLK

Epitope name GGK

Immunogen HIV-1 infection
Species (MHC) human (B8)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius et al. [2000, 2001a] in an IFNgamma elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there
  was no correlation between CTL responses with viral rebound
  rates, plateau viral loads, or clearance rates.

**HXB2 Location** p17 (24–35)

**Author Location** p17 (25–35 SF2)

Epitope GGKKKYKLKHIV

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords review, immunodominance, escape

References Goulder et al. 1997a; Phillips et al. 1991

 Longitudinal study of CTL escape mutants in people with the appropriate HLA types – little variation was observed in the immunodominant B27 epitope, relative to B8 epitopes, which varied over time. • Goulder et al. [1997a] is a review of immune escape that points out that there may be a protective effect associated with B27. and that HLA-B8 individuals tend to progress more rapidly than HLA-B27 patients.

**HXB2 Location** p17 (24–35) Author Location p17 (25-35) Epitope GGKKKYKLKHIV Immunogen HIV-1 infection Species (MHC) human (B8) References Birk et al. 1998b

• A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

**HXB2 Location** p17 (27–35) Author Location p17 (27–35) **Epitope** KRYMIKHLV Subtype C

Immunogen HIV-1 infection Species (MHC) human (Cw\*0602)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

HXB2 Location p17 (28-36) Author Location (C consensus) Epitope HYMLKHIVW Subtype C Immunogen HIV-1 infection Species (MHC) human (A\*2301) Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the M in the third residue HYM-LKHIVW are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** p17 (28–36) **Author Location** (C consensus) **Epitope** HYMLKHLVW

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (A\*2301, A\*2402)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords cross-presentation by different HLA, characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p17 (28–36) Author Location p17 (28-36 LAI) Epitope KYKLKHIVW Subtype B

Immunogen

Species (MHC) human (A\*2402)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes that this is an A\*2402 epitope.

HXB2 Location p17 (28-36) **Author Location** p17 (28–36 SF2) Epitope KYKLKHIVW Immunogen HIV-1 infection Species (MHC) human (A\*2402)

References Ikeda-Moore et al. 1998

- Strong CTL activity to this peptide was detected in 2/3 HIVinfected individuals who were HLA A24+.
- HLA A24 is very common in Japanese (70% carry it) and is common globally.
- · This epitope was detected by looking for peptides with appropriate A24 anchor residues (Y at position 2, carb-term ILF or W) - 16/17 such peptides bound to A24 - KYKLKHIVW was found to be a naturally processed epitope that elicits a strong CTL response.

**HXB2 Location** p17 (28–36) **Author Location** (28–36)

Epitope HYMLKHLVW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

Assay type Other

**Keywords** HLA associated polymorphism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- HYMLKHLVW was a previously defined A\*2402 presented epitope that encompassed an A\*24 associated polymorphism, HYmLKHLVW,in the third position.

HXB2 Location p17 (28-36) Author Location p17 (28–36 LAI) **Epitope** KYKLKHIVW Subtype B Immunogen

Species (MHC) human (A23)

References Goulder & Walker 1999

• P. Goulder, pers. comm.

HXB2 Location p17 (28-36) Author Location p17 (28–36 LAI) **Epitope** KYKLKHIVW Subtype B **Immunogen** Species (MHC) human (A24)

References Brander & Walker 1996

• D. Lewinsohn, pers. comm.

**HXB2 Location** p17 (28–36) **Author Location** p17 (28–36 SF2) Epitope KYKLKHIVW Immunogen HIV-1 infection Species (MHC) human (A24) Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- · Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- · The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A24+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 2/3 group 2, and 0/0 group 3.

**HXB2 Location** p17 (28–36) **Author Location** p17 (28–36 93TH253 subtype CRF01)

Epitope KYKLKHIVW Subtype CRF01\_AE Immunogen HIV-1 infection Species (MHC) human (A24) **Keywords** subtype comparisons References Bond et al. 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- The only HLA-A24 FSWs tested did not recognize the E clade version of this epitope KYKMKHLVW, which differs from the previously defined B clade version by two amino acids, KYKLKHIVW.

**HXB2 Location** p17 (28–36) **Author Location** p17 Epitope KYKLKHIVW

Epitope name KW9

Immunogen HIV-1 infection Species (MHC) human (A24)

**Donor MHC** A2, A24 B38, B60, Cw2, Cw12 Assay type CD8 T-cell Elispot - IFNγ

Keywords supervised treatment interruptions (STI), acute/early infection

References Montefiori et al. 2003

• HIV-1 + patient AC10 underwent early HAART treatment, which was discontinued 1.5 years later. At this timepoint potent NAb responses against autologous virus were detected. Treatment interruption initially induced weak CD8+ responses directed against 5 epitopes. By days 873d and 923d the CTL response had broadened to target 22 epitopes; of these six were fully characterized. Eventually the virus escaped the NAb response, but escape was not accompanied by a rise in viral load, and the authors suggest the virus was contained by the CTL response.

HXB2 Location p17 (28–36) Author Location p17 (28-36) Epitope KYKLKHIVW Immunogen HIV-1 infection Species (MHC) human (A24)

**Donor MHC** A\*0201, A\*2402, B\*52, B75, Cw\*03; A\*0207, A\*2402, B\*46, B\*52, Cw\*01; A\*2402, A\*26, B\*07, B\*5101, Cw\*07

Country Japan.

Assay type Chromium-release assay Keywords epitope processing, escape References Yokomaku et al. 2004

- Epitope variants escaped from being killed by CTLs in an endogenous expression system although they were recognized when corresponding synthetic peptides were exogenously loaded onto the cells. Escape is thus probably due to changes that occur during the processing and the presentation of epitopes in infected cells.
- Epitope variants recognized when added exogenously but not when processed endogenously were: kyRlkhLvw, RyRlkhLvw and QyRlkhivw.

**HXB2 Location** p17 (28–36)

**Author Location** p17 (28–36)

Epitope KYKLKHIVW

Epitope name QW9

Immunogen HIV-1 infection

Species (MHC) human (A24)

**Donor MHC** A\*24, A\*30, B\*39, B\*47, Cw\*12, Cw\*17; A\*24, A\*23, B\*39, B\*07, Cw\*12, Cw\*17

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection, reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs.
   Escape variants were commonly detected in maternal plasma.
   Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- qYKLKHIVW is an escape variant of the A\*24 epitope KYK-LKHIVW, found in 9/10 clones from the mother. It was transmitted to her infant, and persisted for 15 months. Both the mother and child are A\*24+.
- qYKLKHIVW elicted lower responder cell frequencies than KYKLKHIVW.

**HXB2 Location** p17 (28–36)

**Author Location** p17 (728–736 subtype A)

Epitope KYRLKHLVW

Subtype A

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (Cw4)

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-Cw4 women, 2/2 HEPS and 7/11 HIV-1 infected women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in both of the 2/2 HEPS cases and in 3 of the 7/11 HIV-1 infected women.

HXB2 Location p17 (28-36)

**Author Location** p17 (28–36)

Epitope KYRLKHLVW

Immunogen HIV-1 infection

Species (MHC) human (Cw4)

References Appay et al. 2000

- This epitope is newly defined in this study.
- Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virusspecific CD8+ T cells produced IFN-γ and MIP-1β with a distinct subset that failed to produce TNF-α.

**HXB2 Location** p17 (28–36)

**Author Location** 

Epitope KYRLKHLVW

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls (ML1573).

**HXB2 Location** p17 (33–41)

Author Location p17 (33-41)

Epitope HLVWASREL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0602)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

- PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.
- This is a novel predicted epitope.

**HXB2 Location** p17 (33–41)

**Author Location** p17

**Epitope** HLVWASREL

Epitope name HL-9

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0804)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Eli-

spot - IFN $\gamma$ , Intracellular cytokine staining,

Chromium-release assay

Keywords subtype comparisons, epitope processing, im-

munodominance, cross-presentation by different HLA

References Masemola et al. 2004b

- Highly targeted regions in Gag for CD8+ T-cells were defined for individuals with C clade infections in South Africa. 9 specific epitopes within the most reactive regions were characterized. This is 1 of 5 novel epitopes that were found among subtype C HIV-1 from African patients who hadn't previously been identified in B clade infections. Some epitopes were shown to be promiscuous, presented by multiple class I restricting alleles.
- HLVWASREL was presented by Cw\*08 and newly identified in this study; Cw\*08 is slightly more common in Zulus than Caucasians (0.066 versus 0.038).

HXB2 Location p17 (33-41)

**Author Location** 

**Epitope HLVWASREL** 

**Immunogen** 

Species (MHC) human (Cw\*0804)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes that this is an Cw\*0804 epitope.

**HXB2 Location** p17 (34–44)

**Author Location** p17

Epitope LVWASRELERF

Epitope name LF-11

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (A\*3002, B\*570301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Eli-

spot - IFN $\gamma$ , Intracellular cytokine staining,

Chromium-release assay

**Keywords** subtype comparisons, epitope processing, immunodominance, cross-presentation by differ-

ent HLA

References Masemola et al. 2004b

 Highly targeted regions in Gag for CD8+ T-cells were defined for individuals with C clade infections in South Africa. 9 specific epitopes within the most reactive regions were characterized. This is 1 of 5 novel epitopes that were found among subtype C HIV-1 from African patients who hadn't previously been identified in B clade infections. Some epitopes were shown to be promiscuous, presented by multiple class I restricting alleles. • LVWASRELERF was clearly presented by both A\*3002 and B\*570301, it might also be cross-presented by A\*3001, but not as effectively. A\*30 is 10-fold more common among Zulus than Caucasians (allele frequency 0.195 versus 0.019), while B\*57 is similar (0.051 versus 0.043).

HXB2 Location p17 (34-44)

Author Location p17 (34-44)

Epitope LVWRELERF

Immunogen

Species (MHC) human (A30)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an A30 epitope.

**HXB2 Location** p17 (34–44)

**Author Location** (C consensus)

Epitope LVWASRELERF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5703)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- LVWASRELERF is an optimal epitope.

**HXB2 Location** p17 (34–44)

**Author Location** (C consensus)

**Epitope** LVWASRELERF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p17 (36–44)

Author Location p17 (35-43 LAI)

Epitope WASRELERF

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B\*3501)

References Goulder et al. 1997d

- Optimal epitope defined from within p17(30-44), LKHIVWAS-RELERFA.
- Dominant CTL response in an HIV+ asymptomatic donor was to this epitope.
- The Phe in the C-term anchor is distinct from the previouslydefined Tyr for B\*3501 C-term anchors.

**HXB2 Location** p17 (36–44)

Author Location p17 (36-44 LAI)

**Epitope WASRELERF** 

Subtype B

Immunogen

Species (MHC) human (B\*3501)

Keywords optimal epitope

References Frahm et al. 2007; Goulder et al. 1997b

• C. Brander notes this is a B\*3501 epitope.

**HXB2 Location** p17 (36–44)

Author Location p17 (36-44)

**Epitope** WASRELERF

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Birk et al. 1998b

 A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

**HXB2 Location** p17 (36–44)

Author Location p17 (36-44)

**Epitope** WASRELERF

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p17 (36–44)

**Author Location** p17 (36–44 SF2)

**Epitope** WASRELERF

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.

• Number of HLA-B35+ individuals that had a CTL response to this epitope broken down by group: 0/2 group 1, 0/2 group 2, and 1/1 group 3.

HXB2 Location p17 (36-44)

**Author Location** 

**Epitope WASRELERF** 

Epitope name Gag-WF9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Sabbaj et al. 2003

Among HIV+ individuals who carried HLA B35, 1/21 (5%) recognized this epitope.

HXB2 Location p17 (36-44)

**Author Location** Gag

Epitope WASRELERF

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** HIV exposed persistently seronegative (HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- No one, 0/3 HLA B35+ infection-resistant men, and 0/5 preseroconversion men who went on to become infected, reacted to this epitope.

**HXB2 Location** p17 (36–44)

Author Location p17 (36-44)

**Epitope** WASRELERF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0602)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

HXB2 Location p17 (36–44) Author Location p17 (SF2) Epitope WASRELERF Immunogen HIV-1 infection Species (MHC) human

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- The dominant response in an African American who was HLA A3/33 B35/B53 Cw4/7 was to this epitope, although the restricting element was not determined – this epitope fell outside the most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

**HXB2 Location** p17 (43–51)

Author Location p17

Epitope RFAVNPGLL

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human (B63)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, cross-presentation by different HLA

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This is a putative HLA-B63 epitope is contained within a reactive peptide containing the B58 supertype binding motif. There is no evidence for B57/B58 cross-presentation of this epitope.

 $\textbf{HXB2 Location} \ \ p17 \ (69\text{--}93)$ 

Author Location p17 (69–93 BH10)

Epitope QTGSEELRSLYNTVATLYCVHQRIE

Immunogen HIV-1 infection Species (MHC) human (A2) References Johnson *et al.* 1991

• Gag CTL response studied in three individuals.

**HXB2 Location** p17 (71–79)

Author Location p17

**Epitope GTEELRSLY** 

Subtype A

Immunogen HIV-1 infection Species (MHC) human (A\*0101)

**Donor MHC** A\*0101, A\*0301, B\*0801; A\*0201, A\*3101,

B\*3501, B\*3905

Country United Kingdom.

**Assay type** CD8 T-cell Elispot - IFNγ, HLA binding **Keywords** escape, acute/early infection, characterizing

CD8+ T cells

References Milicic et al. 2005

- CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient. In an early acute infection of the donor, the precise timing of transmission determines the viral variants transmitted.
- Escape mutation gteelrsIF in this epitope resulted in 98% reduction in HLA binding affinity, and was the transmitted variant.

**HXB2 Location** p17 (71–79)

**Author Location** p17

**Epitope** GSEELRSLY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0101)

**Donor MHC** A\*0101, A\*0301, B\*0801, B\*5101; A\*0101,

B\*0801

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFNγ, HLA binding

**Keywords** escape, acute/early infection, characterizing

CD8+ T cells

References Milicic et al. 2005

- CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. CTL escape variants were often transmitted. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient.
- The second donor in the study shares A\*0101 and B\*0801 with his partner. Escape mutations gseeIKsly in GSEELRSLY resulted in 44% reduction in HLA binding affinity and no response in an Elispot assay, and gseeIKsly was the transmitted form

**HXB2 Location** p17 (71–79)

**Author Location** p17 (71–79 LAI)

**Epitope** GSEELRSLY

Subtype B

Immunogen

Species (MHC) human (A1)

References Brander & Walker 1996

• P. Goulder, pers. comm.

**HXB2 Location** p17 (71–79)

Author Location p17 (71-79)

**Epitope** GSEELRSLY

Immunogen HIV-1 infection Species (MHC) human (A1) References Birk et al. 1998b

• A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

HXB2 Location p17 (71-79) Author Location p17 (71-79 HXB2) **Epitope** GSEELRSLY Epitope name GSE Subtype B Immunogen HIV-1 infection

Species (MHC) human (A1) Keywords HAART, ART, supervised treatment interruptions (STI), immunodominance, acute/early

infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load - three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- This epitope was not recognized by the 6/8 study subjects that were HLA-A1.

**HXB2 Location** p17 (71–79) **Author Location** p17 (71–79) **Epitope** GSEELRSLY

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A1)

Keywords HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-A1 women, 1/1 HEPS and 3/3 HIV-1 infected women recognized this epitope, and the response was the dominant HLA-A1 response in all cases.

HXB2 Location p17 (71-79) Author Location p17 **Epitope GSEELRSLY Epitope name** GSE Immunogen HIV-1 infection Species (MHC) human (A1)

Keywords HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius et al. [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with supervised treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there was no correlation between CTL responses with viral rebound rates, plateau viral loads, or clearance rates.

**HXB2 Location** p17 (71–79) Author Location p17 (71–79) **Epitope** GSEELRSLY Immunogen HIV-1 infection Species (MHC) human (A1) Country Spain.

> Assay type proliferation, CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 3/13 patients recognized this epitope.

**HXB2 Location** p17 (71–79)

**Author Location** (71–79 B consensus)

**Epitope** GSEELRSLY

Epitope name GY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A1)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords characterizing CD8+ T cells

References Allen et al. 2004

- This study characterizes an escape mutation in a C-terminal flanking residue of the HLA-A3 gag p17 KK9 epitope that inhibits processing, and is embedded in the overlapping HLA-A3 RK9 epitope.
- The immune response was tracked in subject AC-38. The acute immunodominant response was to the B57 TW10 epitope; this response declined following viral escape (tsNlqeqigw) by day 64. The p17 KK9 and RK9 became immunodominant, but then declined as the escape mutation arose. Three other strong responses that persisted were detected, along with 1 sub-dominant response to GY9.

**HXB2 Location** p17 (71–79) **Author Location** Gag **Epitope GSEELRSLY** Epitope name GL9 Subtype B Immunogen HIV-1 infection Species (MHC) human (A1)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

 $\textbf{Keywords} \ \ \text{subtype comparisons, escape, characterizing}$ 

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 9, gseelrslF, was found in the most polymorphic residue in the epitope. These were shared between clades B and C.

**HXB2 Location** p17 (71–79)

**Author Location** p17

**Epitope** GSEELRSLY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A1)

**Donor MHC** A1, A1, B8, B55, Cw3, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence had dramatic changes, the epitope GSEELRSLY peptide was GtegikSLh, and so likely not the actual reactive epitope in the larger peptide.

**HXB2 Location** p17 (71–85)

**Author Location** p17 (71–85 SF2)

**Epitope** GSEELRSLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A1, A11, B8, B27.

HXB2 Location p17 (71-85)

**Author Location** p17 (71–85 HXB2)

**Epitope** GSEELRSLYNTVATL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 were chronically infected and treated; 22 started treatment during acute infection; 11 continuously treated and 11 with STI
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- Responses to this peptide were detected in 16% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p17 (71–90)

Author Location Gag (HXB2)

Epitope GSEELRSLYNTVATLYCVHQ

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A2

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** assay standardization/improvement, HAART, ART

References Chitnis et al. 2003

- 17 perinatally HIV-1 infected children (0.08-16 years) were evaluated for HLA-A2-restricted IFN-gamma CD8+ CTL responses against 4 immunodominant peptides that carry HLA-A2 epitopes. Two peptides were from gp120 (one at position 112, one from the V3 loop), and one each was from gp41 and Gag. 15/17 patients responded to the Gag peptide, 13/17 to the gp41 and the non-V3 gp120 peptides, and 11/17 responded to the V3 loop. 4 children recognized all 4 peptides.
- In 10/14 children, addition of exogenous IL-15 induced increased frequencies of SFCs to the Gag peptide. IL-2 and IL-7 did not increase SFCs, however IL-2, IL-7 and IL 15 could all increase the intensity of the spots in some patients. In 4 children, IL-15 addition brought the SFC response up to the level of detection.

**HXB2 Location** p17 (73–82)

Author Location p17 (73–82)

**Epitope** EELRSLYNTV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4006)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

 $\textbf{Keywords} \ \ \text{subtype comparisons, computational epitope}$ 

prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** p17 (74–82)

Author Location p17

**Epitope** ELRSLYNTV

Immunogen

Species (MHC) human (B\*0801)

Keywords optimal epitope

References Frahm et al. 2007

• Noted by Brander to be a B\*0801 epitope.

**HXB2 Location** p17 (74–82)

**Author Location** p17

**Epitope** ELRSLYNTV

Immunogen

Species (MHC) human (B8)

References Goulder et al. 1997g

• Defined in a study of the B8 binding motif.

HXB2 Location p17 (74-82)

Author Location p17 (74–82)

Epitope ELRSLYNTV Immunogen HIV-1 infection

Species (MHC) human (B8)

References Birk et al. 1998b

• A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

HXB2 Location p17 (74-82)

**Author Location** p17 (74–82)

Epitope ELRSLYNTV

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location p17 (74-82)

**Author Location** p17 (74–82)

**Epitope** ELRSLYNTV

Immunogen HIV-1 infection Species (MHC) human (B8) References Day et al. 2001

• B8-restricted CTL accounted for about 1/3 of the total CTL response in one individual.

**HXB2 Location** p17 (74–82)

**Author Location** (B consensus)

**Epitope** ELRSLYNTV

Epitope name EV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A11, A29, B08, B44, Cw4, Cw7

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

 $\textbf{Keywords} \ \ \text{assay standardization/improvement, memory}$ 

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** p17 (74–82)

**Author Location** p17

**Epitope** ELRSLYNTV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A1, A1, B8, B55, Cw3, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence ELRSLYNTV had dramatic changes, the epitope peptide was gikSLhNTV, and so likely not the actual reactive epitope in the larger peptide.

**HXB2 Location** p17 (74–83)

**Author Location** Gag

**Epitope** ELRSLYNTVA

Epitope name 1241

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for ELRSLYNTVA: 71%. This epitope was previously identified in the literature, but was not confirmed in this study.

**HXB2 Location** p17 (76–86)

Author Location p17 (74–86 LAI)

**Epitope** RSLYNTVATLY

Subtype B

Immunogen

Species (MHC) human (A\*3002)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*3002 epitope.

**HXB2 Location** p17 (76–86)

Author Location p17 (SF2)

Epitope RSLYNTVATLY

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in a single HIV+ individual from Boston this epitope fell outside the most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

**HXB2 Location** p17 (76–86)

Author Location Gag (96ZM651.8)

**Epitope** RLSYNTVATLY

Immunogen

Species (MHC) human (A\*3002)

References Novitsky et al. 2001

- This study provides a survey of CTL responses and full length HIV-1 genome sequences from a C subtype infected Botswanan cohort.
- Only 3/13 (23.1%) A\*3002-positive subjects demonstrated moderate CTL responses to the peptide GTEELRSLYNTVAT-LYCVHE (residues 71 to 90), which contains the previously described A\*3002 epitope RLSYNTVATLY.

**HXB2 Location** p17 (76–86)

**Author Location** p17 (76–86)

**Epitope** RSLYNTVATLY

Epitope name RY11 (p17)

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

References Goulder et al. 2001a

- HLA-A\*3002 is very common in African populations, 50% of Zimbabweans express HLA-A30, 44% in African Zulu, so five new HIV epitopes were characterized that are presented by this HLA molecule.
- A rapid method was developed combining ELISPOT with intracellular IFN-γ staining of PBMCs to map optimal epitopes, then HLA presenting molecules were defined this method was completed within 48 to 72 hours of receipt of blood.
- Two individuals were studied: Subject 199 (HLA A\*0201/\*3002 B\*4402/51 Cw2/5), a Caucasian, and Subject 6007 (HLA A\*3002/ B53/\*5801 Cw4/7) an African-Caribbean.
- In both HLA-A\*3002 individuals the response to RSLYNT-VATLY was dominant.
- Three quantitative assays, ELISPOT, precursor frequency and chromium release, confirmed a hierarchy of response: RY11 (p17) > KY9 (gp41) > KY9 (RT-53) > IY9 (gp41).
- HLA-A\*3001-positive targets do not present RSLYNTVATLY.

**HXB2 Location** p17 (76–86)

**Author Location** 

Epitope RSLYNTVATLY

Epitope name Gag-RY11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

**Donor MHC** A\*3002 A\*3201 B\*4501 B\*5301 Cw\*0401

Cw\*1202

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- Subject 00RCH33 was on HAART had a viral load of 2900 and CD4 count of 727 and also recognized the epitopes YPLTFG-WCY, Nef(135-143), HLA B\*5301; AETFYVDGA, RT(437-445), HLA B\*4501; and HIGPGRAFY, gp160(310-318), HLA A\*3002.

 Among HIV+ individuals who carried HLA B30, 3/16 (19%) recognized this epitope.

**HXB2 Location** p17 (76–86) **Author Location** (C consensus)

Epitope RSLYNTVATLY

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*3002) Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$  Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p17 (76–86)

Author Location (C consensus)

Epitope RSLYNTVATLY

Subtype C

**Immunogen** HIV-1 infection **Species (MHC)** human (A\*3002)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- RSLYNTVATLY is an optimal epitope.

**HXB2 Location** p17 (76–86)

Author Location p17 (74–86 SF2)

**Epitope** RSLYNTVATLY **Immunogen** HIV-1 infection

Species (MHC) human (A30)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic

infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.

- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A30+ individuals that had a CTL response to this epitope broken down by group: 0/1 group 1, 0/0 group 2, and 1/1 group 3.

HXB2 Location p17 (76-86)

Author Location p17

Epitope RSLYNTVATLY

**Epitope name** A30-RY11(p17)

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A30)

**Donor MHC** A30, A32, B18, B27

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8+ T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals.
   Patient D displayed the greatest response to B27-KK10 (p24), and also responded to A30-RY11(p17), A32-PW10(RT), A30-KY11(RT), A32-RW10(gp120), and B18-YY9(Nef).

HXB2 Location p17 (76-86)

**Author Location** p17

Epitope RSLYNTVATLY

Epitope name RY-11

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A30)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining,

Chromium-release assay

**Keywords** subtype comparisons, epitope processing, immunodominance, cross-presentation by differ-

ent HLA

References Masemola et al. 2004b

- Highly targeted regions in Gag for CD8+ T-cells were defined for individuals with C clade infections in South Africa. 9 specific epitopes within the most reactive regions were characterized.
- RSLYNTVATLY was presented by A\*30, which is more common in Zulus than Caucasians (0.195 versus 0.019). This epitope had previously identified in B clade infections.

HXB2 Location p17 (76-86)

Author Location p17

**Epitope** RSLYNTATLY

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A30)

**Donor MHC** A\*02, A\*30, B\*4402, B\*15

Assay type T-cell Elispot

**Keywords** HIV exposed persistently seronegative (HEPS)

References Missale et al. 2004

- HIV-specific T-cell response was tested in patients exposed to blood from a patient with highly replicating HIV; these patients were nosocomially infected with HBV, but uninfected with HIV. HIV-specific T-cell responses were directed to structural and non-structural HIV proteins in 2 patients, suggesting that the virus replicated in these patients sufficiently to prime a cell-mediated immune response that protected them from HIV infection
- This patient responded to 4/8 HIV epitopes tested in an IFNgamma Elispot assay or tetramer assay. Responses were detected 8 and 28 weeks after exposure, this particular epitope was only tested with Elispot.

**HXB2 Location** p17 (76–86)

**Author Location** 

**Epitope** RSLYNTVATLY

Immunogen

Species (MHC) human (B58)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes that this is an B58 epitope.

**HXB2 Location** p17 (76–86)

**Author Location** 

**Epitope** RSLYNTVATLY

Immunogen

Species (MHC) human (B63)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes that this is an B63 epitope.

**HXB2 Location** p17 (76–86)

Author Location p17

**Epitope** RSLYNTVATLY

Epitope name RY11

Subtype B, C

Immunogen HIV-1 infection

**Species (MHC)** human (B63, B57, B58)

**Donor MHC** A\*02, A\*24, B\*1517, B\*58, Cw\*03, Cw\*07

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, cross-presentation by dif-

ferent HLA, optimal epitope

## References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This is a putative HLA-B63/57/58 epitope containing the B58 supertype binding motif. Peptide reactivity was enriched for those that carry B63, with a trend for those that carry B57/B58. Optimal epitope was defined in an individual that was B\*1517(B63)/B58 positive.

HXB2 Location p17 (77-85)

Author Location p17 (77-85)

Epitope SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A\*02)

Keywords HAART, ART

References Huang et al. 2000

- The single cell ELISPOT assay was optimized and highly specific, and found to work well even after the primary cells had been frozen and thawed.
- Increases in gamma IFN producing cells were observed in response to anti-retroviral therapy using single cell IFN-gamma-production ELISPOT.
- 4/8 A\*02 subjects had a positive response to this epitope indicating that it is a major epitope for CD8+ gamma IFN production.
- In 3/3 HLA A\*02, B\*27 individuals, the dominant response in gag measured by both gamma IFN production and T-cell lysis was a B27 epitope, p24(263-272), not the A2 SLYNTVATL epitope.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85)

Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection

**Species (MHC)** human (A\*02)

Keywords HAART, ART

References Rinaldo et al. 2000

Administration of triple-drug antiretroviral therapy (IDV, 3TC and ZDV) sometimes showed a transient increase and other times failed to increase CTL responses in patients with advanced HIV disease, but there is a stable population of tetramer stained HIV-specific CD8+ CD45RO+ cells that can persist after therapy and long periods of virus being below the level of detection.

**HXB2 Location** p17 (77–85)

Author Location p17

Epitope SLYNTVATL

**Epitope name** SL9

Immunogen HIV-1 infection

Species (MHC) human (A\*02)

Keywords HAART, ART, immunodominance

References Scott-Algara et al. 2001

- This study examined with CTL response in HLA A\*02+ children by tetramer staining for HLA-A2 immunodominant epitopes SLYNTVATL and ILKEPVHGV.
- 71% of the 28 HIV-1 infected HLA-A\*02 positive children recognized both epitopes, with cells from 26 children stained positive by the gag tetramer (SLYNTVATL) and 21 children by the pol tetramer (ILKEPVHGV)
- There were no differences observed in children that had therapy versus those that did not.
- Tetramer-binding cells were memory activated CD28-, CD45RO+, CD45RA- HLADR+, CD69-, CD8+ T-cells.

HXB2 Location p17 (77-85)

Author Location p17

**Epitope** SLYNTVATL

**Epitope name** GAG

Immunogen HIV-1 infection

Species (MHC) human (A\*02)

Country France.

**Assay type** Cytokine production, Tetramer binding, Intracellular cytokine staining, Flow cytometric

T-cell cytokine assay **Keywords** responses in children, characterizing CD8+ T

cells

References Scott-Algara et al. 2005

 Only a fraction of HIV-1-specific CD8 T-cells detected in the PBMC of 17 infected children (ages 2-18) were able to produce cytokines (IFN-gamma, TNF-alpha) or chemokines (CCL4, CCL5) after stimulation with the cognate peptide. A negative correlation was found between the plasma viral load and the precentage of CD8+ Gag-specific T-cells secreting IFN-gamma. Tetramers used in this study were SLYNTVATL-HLA-A\*02 and ILKEPVHGV-HLA-A\*02.

HXB2 Location p17 (77-85)

**Author Location** p17 (77–85 HXB2)

Epitope SLYNTVATL

Epitope name SL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords epitope processing, immunodominance, es-

cape

References Brander et al. 1999

- Multiple natural variations in the SL9 flanking regions of the immunodominant epitope SLYNTVATL were tested and found not to adversely affect CTL recognition or prevent epitope processing, suggesting that viral escape from the HLA-A\*0201restricted CTL response against SLYNTVATL is probably not linked to variations in the flanking regions of this epitope.
- The substitution Y79F was an escape mutation in that it interfered with CTL recognition by one CTL clone from an A\*0201 infected individual, clone 13010.B17, but it was still recognized by another CTL clone, 115.D4.

**HXB2 Location** p17 (77–85)

**Author Location** p17

**Epitope** SLYNTVATL **Immunogen** HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** acute/early infection

References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T-cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects – 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

**HXB2 Location** p17 (77–85)

**Author Location** Gag

**Epitope SLYNTVATL** 

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Tan et al. 1999

Adoptive transfer of two autologous in vitro-expanded CTL clones against the A\*0201 restricted epitopes SLYNTVATL and VIYQYMDDL were infused into a patient – they were well tolerated, but the SLYNTVATL clone was shown by tetramer staining to be rapidly eliminated through apoptosis, and the treatment had no impact upon viral load and CD4 and CD8 cell counts.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85)

Epitope SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant
- 95 optimally-defined peptides from this database were used to screen for INFγ responses to other epitopes.
- Individuals who did not respond to SLYNTVATL recognized other HIV epitopes, and 2/4 SLYNTVATL responders had stronger responses to epitopes restricted by other class I alleles.
- SLYNTVATL was the only response detected in a one individual that was HLA A\*0201, B44, B70.

HXB2 Location p17 (77–85)

**Author Location** p17 (77–85) Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A\*0201) Keywords HAART, ART

References Ogg et al. 1999

- CTL effector levels were measured after potent ARV therapy using HLA-tetramer complexes for the A\*0201 epitopes SLYNTVATL and ILKEPVHGV in seven patients, and the B\*3501 epitope DPNPQEVVL in one additional patient.
- Levels of CTL effectors typically decline for 5-7 days and then rebound, fluctuating during the first two weeks of therapy.
- After the early fluctuation, there was a steady exponential decay with a median half-life of 45 days.

HXB2 Location p17 (77-85) Author Location p17 (77–85)

**Epitope SLYNTVATL** 

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

References Altman et al. 1996

- This paper introduces the tetramer methodology that permits quantification of specific CTL based on expression of specific TCRs - HLA-A2 tetramers were prepared that can stain CTL lines specific for ILKEPVHGV and SLYNTVATL, and quantitate HIV-specific CD8+ cell lines in freshly isolated PBMCs.
- Three patients only stained the Gag epitope SLYNTVATL, one patient had the highest frequency of tetramer staining to the Pol epitope (0.77%), less to the Gag epitope (0.28%).

**HXB2 Location** p17 (77–85) **Author Location** Gag

Epitope SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Keywords HAART, ART References Gray et al. 1999

• Administration of highly active antiretroviral therapy (HAART) reduced CD8+ cell frequency, and the CD8+ cells detected by tetramer staining were likely to be memory cells, indicating that persistently replicating viral populations are needed to maintain high frequencies of HIV-1 specific CTL.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85 SF2)

**Epitope SLYNTVATL** 

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** subtype comparisons

References McAdam et al. 1998

• CTL from a patient infected with clade B virus did not recognize the clade A analog of this epitope.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85)

**Epitope** SLYNTVATL

**Epitope name** SL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords TCR usage

References Wilson et al. 1998a

- HIV+ individuals were followed longitudinally using MHC tetramers in combination with 14 anti-BV chain MAbs, and clonal expansion of HIV-specific T cells was followed in vivo.
- Seven HIV+ people were studied, and all showed expansions of particular TCR BV clones, often several, relative to uninfected controls.
- Three patients were followed in detail, TCR VB expansions persisted for 2 to 3 years, with occasional transient increases.
- An A2-Gag specific line from one patient was found to be BV8, and at its highest level represented 17.5% of the patient's CD8+ T cells.

**HXB2 Location** p17 (77–85)

Author Location p17 (77-85)

**Epitope SLYNTVATL** 

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Ogg et al. 1998b

- HLA-tetrameric complexes were used in a cross-sectional study of 14 untreated HLA A\*0201 positive individuals, revealing an inverse relationship between HIV Gag and Pol specific CTL effector cells (CTLe) and viral load.
- Inclusion of both the p17 SLYNTVATL and RT ILKEPVHGV epitopes gives a good representation of HLA A\*0201-restricted activity.
- No correlation was observed between the CTLe and CD4 count or clearance rate of productively infected cells.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85)

Epitope SLYNTVATL

Epitope name SL9

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Keywords epitope processing

References Walter et al. 1997

- HLA-A2 heavy chain and  $\beta$ 2-microglobulin expressed in E. coli were refolded in the presence of this peptide.
- The HLA-A2-peptide complex elicited HLA-A2 peptidespecific CTL response in cells lacking HLA-A2.
- Suggests that preformed HLA-peptide complexes could provide an alternate to intracellular processing for immunogens.

**HXB2 Location** p17 (77–85)

Author Location p17 (77-85)

**Epitope SLYNTVATL** 

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Lalvani et al. 1997

• A peptide-based protocol was optimized for restimulation of CTLp using optimized peptide and IL-7 concentrations - importantly this protocol does not stimulate a primary response, only secondary - peptide-specific CTLp counts could be obtained via staining with peptide-Class I tetramers.

 This peptide was one of the test peptides for optimizing the protocol.

HXB2 Location p17 (77–85) Author Location p17 (76–84) Epitope SLYNTVATL

Epitope name SL9

Immunogen in vitro stimulation or selection

**Species (MHC)** human (A\*0201) **References** van der Burg *et al.* 1996

- Slow dissociation rate is associated with immunogenicity.
- CTL generated by in vitro stimulation of PBMC derived from uninfected individual.

HXB2 Location p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A\*0201) Keywords review, escape

References Goulder et al. 1997e; Goulder et al. 1997a

- HLA-identical siblings, hemophiliac brothers, were both infected with the same batch of factor VIII.
- One had a response to gag A2 epitope SLYNTVATL, the other to pol A2 epitope ILKEPVHGV. They were tested 6-8 years after infection.
- Viral sequencing from the twin that had no response to SLYNT-VATL indicated his virus had the substituted form SLH-NAVAVL.
- 71% of an additional set of 22 HIV-1 infected HLA-A\*0201 positive donors preferentially responded to gag SLYNTVATL.
- Those individuals with a pol ILKEPVHGV response tended to have mutations in or around SLYNTVATL.
- An additional subject went from SLYNTVATL responder to non-responder coincident with a switch to the variant SLFNT-VATL.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study.

**HXB2 Location** p17 (77–85) **Author Location** p17 (77–85)

**Epitope** SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Keywords review

References Goulder et al. 1997a

- This paper is a review of CTL and immune evasion, but it presents a study of a shift from an HLA-A\*0201 response to SLYNTVATL, to a B62 response to GLNKIVRMY.
- As long as a strong CTL response to SLYNTVATL was evident, the epitope variants SLFNTVATL or SLYNTIATL dominated the viral population – eventually the CTL response to the index peptide became undetectable, the CTL response shifted to a focus on GLNKIVRMY, and the index peptide SLYNTVATL once again established itself as the dominant form.

HXB2 Location p17 (77–85) Author Location Gag (77–85) Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords HAART, ART

References Gray et al. 1999

- Peptide-tetramer complexes of A\*0201 and SLYNTVATL or ILKEPVHGV were used to study individuals receiving HAART to determine the frequency of Class I HLA-restricted anti-HIV CD8+ T cells.
- 17/18 asymptomatic patients had a CTL response to one or both epitopes 72% had a CTL response to SLYNTVATL.
- After HAART, the majority of the epitope-specific CTL were apparently memory cells.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85 subtype A)

Epitope SLFNTVATL

**Epitope name** SL9

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** subtype comparisons

References Dorrell et al. 1999

- CTL responses in three individuals with non-clade B infections were studied, two with subtype A infections, one with subtype C – their infections all originated in East Africa.
- This epitope is most commonly SLYNTVATL in B subtype, and CTL from the C subtype infection did not recognize B clade gag or the 3Y form of the epitope, but did recognize the predominant A and C clade form, SLFNTVATL.

**HXB2 Location** p17 (77–85)

Author Location p17 (77-85)

**Epitope** SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** immunodominance

References Brander et al. 1998a

- Of 17 infected HLA A\*0201 subjects, 13 had CTL responses against the p17 SLYNTVATL epitope, six recognized ILKEPVHGV and five recognized VIYQYMDDL, and there was no correlation between viral load and recognition of a specific epitope.
- Only one subject had CTL against all three epitopes.
- There was significant heterogeneity in the CTL response to this immunodominant epitope.
- The overall variation in this epitope among the 17 who had a CTL response and 11 non-HLA A\*0201 HIV-1 + individuals was similar, suggesting a lack of immune pressure.
- Subjects were part of the San Francisco City Clinic Cohort, the ARIEL project and from the Boston area.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85 HXB2)

Epitope SLYNTVATL

Epitope name SL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** rate of progression, immunodominance References Hay et al. 1999

- CTL response to IPRRIRQGL was the immunodominant response in a rapid progressor - there was a subdominant response to SPAIFQSSM in Pol, and interestingly, no response to commonly immunodominant HLA A\*0201 epitope SLYNT-VATL, although this individual was HLA A\*0201.
- The individual showed a strong initial CTL response at the time of the initial drop in viremia, but it was quickly lost, although memory cells persisted.
- Despite the initial narrow response to two epitopes, no other CTL responses developed.
- No HIV-specific lymphoproliferative responses were detected in this patient, and neutralizing antibody response was weak.
- A variant of this epitope was observed in vivo (-F---V-), but this mutation is recognized by SLYNTVATL-specific CTL, and in this case the patient's cells could present the peptide to SLYNTVATL-specific CTL.

HXB2 Location p17 (77-85) Author Location p17 (77–85) **Epitope SLYNTVATL** Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords HAART, ART

References Kalams et al. 1999b

- Two patients were followed before and after HAART reduced plasma HIV-1 RNA levels resulted in a decline in HIV-specific in-vivo activated CTL such that by day 260 CTL activities were undetectable.
- · ERYLKDQQL was the dominant response in one of the individuals, SLYNTVATL subdominant.
- Sporadic breakthrough in viremia resulted in transient increases in CTLp.
- Memory CTL frequency directed against Vac-Gag, Vac-RT, Vac-Env, and Vac-Nef initially increased with HAART and then decreased with the decline of the viral load.

**HXB2 Location** p17 (77–85) **Author Location** Gag (77–85)

**Epitope** SLYNTVATL Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Spiegel et al. 2000

- High levels of CD8+ HIV-1 specific and cytomegalovirus specific CTL were detected by HLA-A\*0201-peptide tetramers in 3 infected subjects with very low CD4 counts, but CD8 T cell mediated effector activity was not seen.
- Thus HIV-1 specific CD8+ cells may be present but may lack direct effector activity in late disease, suggesting that overcoming antigen unresponsiveness may be a useful therapeutic strategy.

HXB2 Location p17 (77-85)

**Author Location** Gag (77–85)

**Epitope** SLYNTVATL

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

References Larsson et al. 1999

- ELISPOT was used to assay the CD8+ T-cell response to the HIV-1 proteins Gag, Pol, Nef or Env expressed in vaccinia vectors in 19 HIV+ people.
- In A\*0201 individuals, higher numbers of spot-forming T cells were directed against HIV-1 proteins expressed in vaccinia than to peptides SLYNTVATL and ILKEPVHGV presented by A2.

**HXB2 Location** p17 (77–85)

Author Location p17 (SF2)

**Epitope SLYNTVATL** 

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords subtype comparisons, immunodominance References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in 11/25 HLA A2 (A\*0201 or A\*0202) HIV+ individuals from Boston and in 1/8 HLA A2 HIV+ individuals from Durban.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85 LAI)

**Epitope SLYNTVATL** 

Subtype B

**Immunogen** 

Species (MHC) human (A\*0201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0201 epitope.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85 SF2)

Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords escape, acute/early infection

References Goulder et al. 2001a

- This epitope is targeted by 75% of HLA-A\*0201, HIV+ adults, and the magnitude of the response is inversely correlated with viral load.
- CTL responses to SL9 and autologous SL9 variants were not detected in 11 HLA-A\*0201 positive subjects during acute infection.
- Longitudinal studies of two individuals (AC13 and PI004) showed that the initial control of viremia was independent of the SL9 CTL response.
- Low Gag expression levels did not correlate with the delayed CTL response to this epitope.

• Autologous SL9 variants SLYNTIAVL, SLYNTVAVL, SLFNT-VATL, SLFNTVATL, and SLFNTVATL are each capable of inducing a range of CTL responses, sometimes strong, sometimes diminished, and sometimes complete escape relative to the wild type variant SLYNTVATL in patients with chronic HIV-1 infection – the ability to cross-react with a particular variant was patient dependent.

**HXB2 Location** p17 (77–85) **Author Location** p17

Epitope SLYNTVATL

**Epitope name** p17 SL9 **Immunogen** HIV-1 infection

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Keywords subtype comparisons, supertype, computa-

tional epitope prediction

References Altfeld et al. 2001c

- HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.
- Three additional previously described HLA-A2 epitopes were added to the set of 20, including p17 SL9, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2).
- p17 SL9 was recognized in 12/22 patients with chronic HIV-1 infection.
- Only 1/13 patients with acute HIV-1 infection recognized p17 SL9.

HXB2 Location p17 (77-85)

**Author Location** Gag

Epitope SLYNTVATL

Epitope name (SL9)

Immunogen HIV-1 infection Species (MHC) human (A\*0201) References Goepfert *et al.* 2000

- This paper describes a comparison of results of different CTL assays, a SL9 tetramer assay and IFN-gamma ELISPOT, using 7 HIV-positive patients.
- The IFN-gamma ELISPOT assay was compared using the single SL9, a pool of overlapping 20 mers, and recombinant vaccinia encoding Gag as antigen – pooled peptides gave the highest number of spot forming cells, vaccinia gave high background.
- A correlation with results of the tetramer assay was found only for ELISPOT using the Gag epitope as antigen, but the tetramer assay detected a 10-fold higher number of cells than could produce IFN-gamma in the ELISPOT assay – the authors suggest not all tetramer-positive cells may produce IFN-gamma, some may be undergoing apoptosis, some may be producing other cytokines.
- The tetramer assay could detect a reaction to SLYNTVATL in most of the HLA-A\*0201 chronically HIV-1 infected study subjects.

**HXB2 Location** p17 (77–85)

**Author Location** Gag (77–85)

**Epitope** SLYNTVATL

Immunogen

Species (MHC) human (A\*0201)

Keywords binding affinity

References Sandberg et al. 2000

 This epitope served as a positive control in a study comparing peptide binding affinity to HLA-A201 to CTL responses upon vaccination with a nef DNA vaccine.

**HXB2 Location** p17 (77–85)

**Author Location** Gag (LAI)

**Epitope** SLYNTVATL

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Keywords dendritic cells

References Engelmayer et al. 2001

- Recombinant canarypox virus vector containing HIV-1 sequences, upon infection of mature dendritic cells, can trigger specific lysis through in vitro by T-cells from HIV-1 infected individuals at levels comparable to the response seen to HIV carried in vaccinia vectors.
- Recombinant canarypox virus vector containing HIV-1 sequences can also stimulate HIV-specific CD4+ helper T-cell responses.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85 LAI)

**Epitope** SLYNTVATL

Epitope name G3

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords HAART, ART

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using tetramer staining or CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+PBL but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

**HXB2 Location** p17 (77–85)

**Author Location** Gag

Epitope SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Gea-Banacloche et al. 2000

 In a study including many long-term non-progressors, no correlation between plasma virus levels and number of HIV-specific CD8+ T-cells was found.

- High frequencies of circulating CD8+ T-cells were HIV-1 specific, and the majority of these responses were to gag-pol gene products.
- 4/21 subjects were HLA-(A\*0201), and of these only 2 subjects (patient 3 and 19) tested positive to this epitope.

HXB2 Location p17 (77–85) Author Location p17 (77–85 SF2) Epitope SLYNTVATL

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- Tetramer staining with A2, beta2microglobulin, and either SLYNTVATL, KLVGKLNWA, or LTFGWCFKL revealed that tetramers detected more HIV-specific sells in LTNP than in progressors, activated effector cells were the minority population, and ELISPOT correlated better with the effector cell subpopulation than the total tetramer stained population.

**HXB2 Location** p17 (77–85)

Author Location Gag (77–85)

**Epitope** SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords HAART, ART, rate of progression

References Jin et al. 2000a

- The CTL effector levels (CTLe) were compared in long term non-progressors (LTNP) with low viral load and in patients whose virus was well-suppressed by therapy, using a tetramer assay.
- LTNPs have high memory CTLe numbers and low viral load, while HAART patients had low CTLe numbers and low viral load.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85)

**Epitope** SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Appay et al. 2000

- Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.

 In most donors, between 50% and 95% of the activated virusspecific CD8+ T cells produced IFN-γ and MIP-1β with a distinct subset that failed to produce TNF-α.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85)

**Epitope SLYNTVATL** 

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Goulder et al. 2000b

- Tetramer assays were compared with three functional assays in 42 people with chronic HIV infection: ELISPOT, intracellular cytokine staining, and precursor frequency (limiting dilution assay [LDA]).
- HIV-specific tetramer staining CTLs appeared to be active, and inert CTL were not found to play a significant role in chronic pediatric or adult HIV infection.

**HXB2 Location** p17 (77–85)

Author Location p17

Epitope SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords dendritic cells

References Ostrowski et al. 2000

- The role of CD4+ T-cell help in expansion of virus-specific memory CTL was studied through co-culture *ex vivo*.
- Optimal expansion of HIV-1-specific memory CTL depended on CD4+ T cell help in 9/10 patients – CD40 ligand trimer (CD40LT) could enhance CTL in the absence of CD4+ T cell help to a variable degree in most of patients.
- Those CTL that didn't respond to CD40LT could expand with IL2 present, and IL15 produced by dendritic cells also contributes
- The T-helper epitope used for CD4+ T cell stimulation was the universal tetanus helper epitope TET830-843 (QYIKANSK-FIGITE).

HXB2 Location p17 (77–85)

**Author Location** 

**Epitope** SLYNTVATL

Subtype B

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost, canarypox prime with gp160 boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Gag, gp120, gp41, Nef, Pol

Species (MHC) human (A\*0201)

Keywords vaccine-specific epitope characteristics

References Ferrari et al. 2001

- Different HIV strains were used for different regions: gp41 LAI, Gag LAI, gp120 MN, gp120 SF2.
- Two vaccinees with Gag responses were HLA-A\*0201+, but neither made SLYNTVATL responses to the Gag vaccine, in contrast to its frequent recognition in natural infections. No HLA-A\*0201 responses were observed to an Env vaccine.

**HXB2 Location** p17 (77–85)

**Author Location** 

**Epitope** SLYNTVATL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Keywords rate of progression, immunodominance

**References** Migueles & Connors 2001

- CTL activity was monitored in 27 individuals, including 10 LTNP with an over-expression of HLA B\*5701 – these individuals have viral loads below the threshold of infection without therapy, and their immune response tends to be focused on peptides that contain B\*5701 epitopes ISPRTLNAW, KAF-SPEVIPMF, TSTLQEQIGW, and QASQEVKNW.
- CTL responses are broader in B\*5701+ individuals with progressive viremia than those that control viremia.
- The HLA-A\*0201 SLYNTVATL epitope response was not as strong in individuals that carried both A2 and B57.

**HXB2 Location** p17 (77–85)

Author Location Gag (77-85)

**Epitope** SLYNTVATL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

 $\label{lem:keywords} \textbf{Keywords} \ \ \text{epitope processing, immunodominance}$ 

References Sewell et al. 2002

- Epitope processing of three different HLA-A\*0201 HIV epitopes was shown to use different pathways, which might influence patterns of immunodominance. 174 cells were used that lack TAP1 and TAP2 genes, as well as the LMP2 and LMP7 genes that encode the beta-subunits of the immunoproteasome. These genes could be added back through transfection to study processing.
- ILKEPVHGV was efficiently presented in TAP-1 and -2 transfected cells while VIYQYMDDL and SLYNTVATL were not. VIYQYMDDL was destroyed by the MB1 subunit of the protease, and could be expressed in the presence of the proteasome inhibitor lactacystin, but SLYNTVATL expression was not restored. SLYNTVATL expression was unaltered by lactacystin in a wild type cell line.

**HXB2 Location** p17 (77–85)

**Author Location** Gag (ADA)

**Epitope SLYNTVATL** 

Epitope name SL-9

Subtype B

Immunogen HIV-1 infected monocyte-derived

Species (MHC) mouse (A\*0201)

References Poluektova et al. 2002

- Nonobese diabetic NOD-C.B-17 SCID mice were reconstituted with HLA-A\*0201 positive human PBL and injected with HIV-1 infected monocyte-derived macrophages MDM in the basal ganglia to provide a mouse model of HIV-1 encephalitis.
- HLA-A\*0201 CTL responses were detected by tetramer staining in the spleen in seven days, increased through day 14, and the numbers of productively infected were reduced >85% in the second week.

HXB2 Location p17 (77–85) Author Location p17 (77–85 LAI)

**Epitope** SLYNTVATL

Epitope name LR23 Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant (IFA), Montanide (ISA 720), P30, PLG

Species (MHC) mouse (A\*0201)

**Keywords** binding affinity, vaccine-specific epitope characteristics, immunodominance

References Peter et al. 2001

- The stability of peptide binding to HLA-A2.1 was determined for six HLA-A2.1 peptides included in this vaccine study

   ILKEPVHGV (RT), SLYNTVATL (p17), SLLNATDIAV (gp41) and LLWKGEGAV (RT) all bound with high affinity comparable to a influenza epitope reference (GILGFVFTL), while RGPGRAFVTI and VIYQYMDDL bound with a lower affinity (relative binding activity = 0.01).
- The four high-affinity peptides formed stable complexes with half-lives ranging between 8 and 32 hours, while the low affinity peptides had half lives of less than an hour.
- HLA-A2.1 transgenic mice were immunized with the six HIV-1 peptides and P30, as a universal T-helper epitope, with IFA or Montanide or microspheres as adjuvants.
- All peptides except VIYQYMDDL induced a stong CTL response in Cr-release assays stronger responses were observed when peptides were delivered alone, indicating immunodominance when the combination was used.

HXB2 Location p17 (77–85)

**Author Location** p17 (77–85 LAI)

**Epitope SLYNTVATL** 

Epitope name LR23

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant

(IFA), IL-12, P30

**Species (MHC)** mouse (A\*0201)

Keywords vaccine-specific epitope characteristics, im-

munodominance

References Peter et al. 2002

• When HIV-1 peptides were used to vaccinate HLA-A2.1 transgenic A2-Kb mice, strong responses to five peptides were observed when the peptides were given individually, but immunodominance limited the response to some of the peptides when they were given in combination Peter *et al.* [2001]. IL-12 can counteract immunodominance in BALB/c mice, so it was given with the multiple epitope vaccination, and was instead found to specifically eliminate the HLA-A2.1-epitope CTL responses, but not Kb CTL responses. This was possibly a consequence of transient depletion of T-cells, B cells and macropahges in the spleen.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85)

**Epitope** SLYNTVATL

Immunogen computer prediction

**Species (MHC)** (A\*0201)

**Keywords** subtype comparisons, computational epitope prediction, vaccine-specific epitope characteristics, escape

References Schönbach et al. 2002

- Computational methods (artificial neural networks [ANN], hidden Markov models [HMM], binding matrices based on HLA association rates BIMAS) were used to identify HLA-A\*0201 and HLA-B\*3501 HIV T-cell epitope candidates from 533 Gag, Env and Pol sequences of which 374 were derived from HIV-1, 97 were derived from HIV-2 and 62 from SIV. Comparisons to known epitopes and between clades were made.
- The SLYNTVATL epitope received focused discussion. SLYNTVATL, slFntvatl, slyntvaVl, and slyntIaVl are all recognized variants, ANN predicts all four variants would be recognized, while BIMAS only predicts SLYNTVATL and slFntvatl would be recognized. However, Sewell et al. [1997] suggested certain substitutions may be antagonistic, including slFntvatl, and vaccines do not stimulate SLYNTVATL responses as well as natural infections. The authors note these kinds of issues complicate the application of computational predictions of epitopes to vaccine design.

HXB2 Location p17 (77–85) Author Location Gag (76–84)

Epitope SLYNTVATL

Subtype B

Immunogen vaccine

Vector/Type: DNA HIV component: HIV-1

Species (MHC) mouse (A\*0201)

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance

References Singh et al. 2002; Sykes & Johnston 1999

- C3H (H-2k) transgenic mice carrying a fused HLA-A\*0201 alpha1 and alpha2 and H-2Dk alpha3 hybrid class I molecule were immunized using an epidermal gene gun with an ubiquitin expression library of 32 plasmids that spanned the HIV-1 genome. Ubiquitin targets the expressed HIV-1 peptides to the proteasome.
- A single immunization with the UB-HIV-1 library vaccine induced potent, stable and multivalent CTL responses against all library members.
- Immunodominant epitopes SLYNTVATL (Gag), ILKEPVHGV (Pol), RIQRGPGRAFVTIGK (Env) and AFHHVAREK (Nef) elicited strong CD8+/IFN- responses and stimulated CTL that were functional in a Cr-release assay and against wild type antigen.
- The presence of multiple plasmids HLA-A\*0201-restricted CTL epitopes did not decrease CTL immunogenicity, and CTL responses to single peptide immunizations were comparable to responses based on mixtures of either 16 or 32 peptides.

HXB2 Location p17 (77-85)

**Author Location** 

**Epitope SLYNTVATL** 

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

**Donor MHC** A\*0202, A\*2501, B\*1801, B62, Cw10,

Cw\*1203, DRB1\*1501, DQB1\*8

Keywords rate of progression, Th1, Th2

References Imami et al. 2002b

- 70 patients with chronic disease progression, 10 clinical non-progressors, and 3 immunologically discordant progressors (individuals who controlled viremia but had progressive CD4+ T-cell decline) were analyzed for their T-helper cell responses to p24 and cytokine profile. Long term non-progressors had much stronger Th responses, particularly to p24 peptides, and they tended to be balanced between Th1, IL-2 producing and Th2, IL-4 producing responses.
  - One of the immunologically discordant progressors became symptomatic during the course of the study, and he had a rapid drop in proliferative response to all antigens and also a shift from a Th1 to a Th2 response. To find out if the CD8 response also shifted in cytokine production, the CD8+ T-cell response to SLYNTVATL in this patient was also tested. It too was found to shift, from IFNgamma to IL-4 producing in Elispot, and using a bioassay of indicator lines, from IL-2 to IL-4 production.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85)

Epitope SLYNTVATL

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

**Donor MHC** A\*0201, A11, B51, B61, Cw2, Cw\*14

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** binding affinity, acute/early infection, early-expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- Only 1/10 HLA A\*02 carrying individuals in this study recognized SLYNTVATL.
- All HIV-1 proteins except Vpu were recognized, and responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

**HXB2 Location** p17 (77–85)

**Author Location** 

**Epitope** SLYNTVATL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Assay type Cytokine production, proliferation, Tetramer binding, Intracellular cytokine staining,

Chromium-release assay

References Dagarag et al. 2003

- Telomer length is short in the CD8+ T-cell compartment of HIV-1 infected people, indicating excessive CTL activation and premature senescense. Here human telomerase RT (hTERT) transduction of HIV-1-specific CTL was used to study the functional impact of telomerase. Telomerase expression enhanced proliferative capacity, as well as cytolytic and antiviral capabilities; cytokine production was unchanged. hTERT transduced CTLs were 10-fold more efficient in controlling HIV-1 replication in culture. Thus telomerase transduction can restore CTL mediated cytolysis, and may have therapeutic potential.
- Three polyclonal CD8+ T-cell lines derived from an HIV-1, HLA A\*0201 positive patient were used in this study, including one specific for this epitope.

HXB2 Location p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL

Subtype B Immunogen vaccine

Vector/Type: peptide HIV component: p24 Gag

Species (MHC) mouse (A\*0201)

Donor MHC A2.1

Assay type Cytokine production, Chromium-release as-

**Keywords** binding affinity, vaccine-induced epitopes **References** Okazaki *et al.* 2003

- Alanine substitutions of VIYQYMDDL were tested for importance of each amino acid for HLA-A2.1 binding. Peptide variant (vLyqymddV) showed an 8 fold higher MHC binding affinity than wild type. YLyqymddV had an even higher binding affinity, but the Y at position one blocked TCR recognition. The higher affinity form of vLyqymddV induced CTL in vivo that could protect against a vaccinia virus expressing RT and the wild type epitope.
- SLYNTVATL was included as a control.

**HXB2 Location** p17 (77–85) **Author Location** Gag (77–85)

**Epitope** SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A\*0201) Assay type Tetramer binding

Keywords genital and mucosal immunity

References Shacklett et al. 2003

- Lymphocytes from rectal biopsies were used to characterize the CD8+ T cell response to HIV in GALT, Gut-associated lymphoid tissues. Patients were selected on the basis of being HLA-A2+ and having detectable SLYNTVATL and ILKEPVHGV tetramer responses in PBMC. SLYNTVATL frequency was increased in GALT relative to PBMC in 6/7 patients studied, while a control response to a CMV-peptide was diminished in GALT. Only two patients had ILKEPVHGV CD8+ T cell responses, and both had slightly higher frequencies in GALT than PBMC.
- HIV may perturb lymphocyte retention in GALT, suggested by an overall reduction of GALT CD8+ cells expressing alphaEbeta7. GALT HIV-specific CD8+ T cells expressed alphaEbeta7, suggesting mucosal priming.

HXB2 Location p17 (77-85)

**Author Location** 

**Epitope SLYNTVATL** 

Epitope name SL9 Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Tetramer binding, T-cell Elispot, Intracellular cytokine staining,

Flow cytometric T-cell cytokine assay

**Keywords** epitope processing, escape, variant cross-recognition or cross-neutralization

References Jamieson et al. 2003

- Epitope escape mutations in chronically infected individuals developed over several years indicating selective advantage of escape mutants. The maturation state of CTLs appear to affect the rate of epitope mutation and CTL decay.
- In two patients, SL9-specific CTL peaked at 2-4 years postinfection; at that point the escape mutations began to dominate followed by CTL decline with a 6 month lag, suggesting CTL decline resulted as a consequence of escape. In a third patient, the initial response was 1/2 as strong and mutations did not arise until 6-7 years post-infection; in that case the decline in SL9 CTL preceded epitope mutation.
- Two patients HLA-A\*0201 started out with a non-consensus sequence, slFntvatl. In one of the patients, a transient reversion to the consensus was observed after 4 years, that did not reappear until the 11th year, suggesting the possibility that a reversion to the consensus form occurred, but a CTL response may have limited it so that this more fit form could not re-assert itself until the patient had a more severely compromised immune response.

HXB2 Location p17 (77-85)

**Author Location** Gag

Epitope SLYNTVATL

Epitope name SL9 Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Country United States.

Assay type Cytokine production, Tetramer binding, Intra-

cellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** epitope processing, rate of progression, immunodominance, acute/early infection, den-

dritic cells, TCR usage, memory cells

References Kan-Mitchell et al. 2004

• SL9-specific CTLs were shown to be primed by immature DCs and independent of help from CD4+ or exogenous IL2, and sensitive to paracrine IL-2 induced apoptosis. The authors suggest that the reason SL9 responses are not seen during acute infection is the high level of innate immune responses resulting in cytokine-induced apoptosis, but that these CD8+ T-cells would come to dominate later infection when CD4 help is diminished.

HXB2 Location p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A\*0201)

**Donor MHC** A\*0201, A\*2402, B\*52, B75, Cw\*03; A\*0201, A\*31, B\*27, B\*5101, Cw\*02; A\*0207, A\*2402, B\*46, B\*52, Cw\*01

Country Japan.

Assay type Chromium-release assay Keywords epitope processing, escape References Yokomaku *et al.* 2004

- Epitope variants escaped from being killed by CTLs in an
  endogenous expression system although they were recognized when corresponding synthetic peptides were exogenously
  loaded onto the cells. Escape is thus probably due to changes
  that occur during the processing and the presentation of epitopes in infected cells.
- Endogenously expressed wild type epitope and slyntIatl variants were recognized by CTL clones while slynLvatl, slFntvaVl and sVyntvatl variants were not. sVyntvatl and slFntvaVl variants were, however, recognized when added exogenously to the cells.

HXB2 Location p17 (77–85) Author Location p17 (77–85 LAI)

**Epitope SLYNTVATL** 

Epitope name SL9 Subtype B

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost, vaccinia prime DNA boost Strain: B clade LAI, B clade MN HIV component: Gag, gp120, gp41, Protease Adjuvant: GM-CSF

Species (MHC) human (A\*0201)

Assay type Cytokine production, CD8 T-cell Elispot -

IFN $\gamma$ , Tetramer binding, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** vaccine-specific epitope characteristics, immunodominance, characterizing CD8+ T cells

References Ferrari et al. 2004

Thirteen HLA-A\*0201 vaccines with anti-Gag CD8+ CTL reactivities were tested in uninfected HIV vaccine recipients to examine the pattern of SL9 epitope immunodominance. None of the vaccines had a detectable anti-SL9 response, in contrast to 75% of HLA A\*0201 chronically infected HIV+ individuals that respond to this epitope.

**HXB2 Location** p17 (77–85)

Author Location p17

**Epitope SLYNTVATL** 

**Epitope name** SL9

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

**Keywords** review, rate of progression, escape, acute/early infection

References Goulder & Watkins 2004

• This paper is a review of the role of CTL in HIV infection, and it uses SL9 as an example of an epitope that is not responded to early in infection, yet 75% of HIV+ people respond to SL9 during chronic infection. Despite the delay in response, strong SL9 responses have been associated with lower viral loads, and escape mutations arise.

HXB2 Location p17 (77-85)

**Author Location** (C consensus)

**Epitope** SLYNTVATL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Country South Africa.

**Assay type** CD8 T-cell Elispot - IFNγ **Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p17 (77–85)

**Author Location** Gag (77–85)

Epitope SLYNTIATL

Epitope name SL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Donor MHC** A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Chromium-release assay

**Keywords** escape, acute/early infection, characterizing

CD8+ T cells

References Bansal et al. 2005

- Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- The response to this peptide was not apparent until month 20, by month 32 a T to V change was dominant, but the slyntiVI mutant showed comparable avidity.

**HXB2 Location** p17 (77–85) **Author Location** Gag (77–85)

Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection, peptide-HLA interaction, vaccine

Vector/Type: peptide Strain: multiple epitope immunogen HIV component: mimotopes Adjuvant: Incomplete Freund's Adjuvant (IFA)

**Species (MHC)** human, transgenic mouse (A\*0201)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, HLA binding

**Keywords** vaccine-specific epitope characteristics, immunodominance, escape, TCR usage, variant cross-recognition or cross-neutralization, vaccine antigen design, mimics

References Boggiano et al. 2005

- A combinatorial library was used to identify epitope mimics of HLA-A2 restricted CTL epitope SL9.
- 19 HIV+ HLA-A\*0201 subjects were tested for their ability to bind to peptide variants. 11/19 could bind to SLYNTVATL. Nine epitope mimics were recognized by more than a third of the subjects, and 1 subject recognized 17/20 variants tested. Some SL9 mimics were up to an order of magnitude better at stimulating CTL responses in PBMC than was SL9.
- Compared to the original SL9 sequence, some SL9 variants recognized by HLA-A\*0201 patients induced superior SL9 immune responses in HLA-A\*0201 transgenic mice.

HXB2 Location p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** assay standardization/improvement, TCR usage, characterizing CD8+ T cells

References Killian et al. 2005

- A novel technique for subtractive analysis of HIV-1 specific CTLs was developed, including depletion of peptide-specific CTLs by stimulating PBMCs with the specific peptide in the presence of 5-FU, followed by TCR spectratyping for clonal breadth analysis. In analysis of infected individuals using this technique, it was found that HIV-1 specific responses range from two to 10 different T-cell clones per epitope.
- The SL9 responses in one individual were complex, with TCR in multiple families, including: Vbeta12.2, Vbeta17, Vbeta23.3 and Vbeta22.
- This paper provides further evidence for the polyclonal nature of epitope-specific responses. Polyclonal responses may be able to better inhibit escape and may play a beneficial role in progression.

HXB2 Location p17 (77-85)

**Author Location** Gag

**Epitope SLYNTVATL** 

Epitope name S9L
Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV compo-

*nent:* gp140, gp140∆V3

**Species (MHC)** transgenic mouse (A\*0201)

Assay type Chromium-release assay, Flow cytometric Tcell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

 A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted CTL responses against multiple conserved HIV-1 epitopes in HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw virus expressing gp140 with deleted V1, V2, and V3 loops successfully induces neutralizing antibodies against HIV-1. A live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination against HIV and measles.

HXB2 Location p17 (77–85) Author Location (C consensus)

Epitope SLYNTVATL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- SLYNTVATL is an optimal epitope.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85)

**Epitope** SLYNTVATL

Immunogen

Species (MHC) human (A\*0202)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

 C. Brander notes that this epitope can be presented by A\*0201 and A\*0202.

**HXB2 Location** p17 (77–85)

Author Location p17 (SF2)

**Epitope** SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A\*0202)

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in 11/25 HLA A2 (A\*0201 or A\*0202) HIV+ individuals from Boston and in 1/8 HLA A2 HIV+ individuals from Durban.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.

• Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), • Sixteen HLA A2+ patients were tested for their ability to make ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

HXB2 Location p17 (77-85) Author Location p17 (77–85 LAI) **Epitope SLYNTVATL** Subtype B Immunogen

Species (MHC) human (A\*0205)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes that this epitope can be presented by A\*0201 and A\*0202.

**HXB2 Location** p17 (77–85) **Author Location** p17 (subtype A) **Epitope SLYNTVATL** Subtype A

**Immunogen** HIV-1 exposed seronegative **Species (MHC)** human (A\*0214, A\*0201)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix – systemic CD8+ T cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.
- The epitope variants SLYNTVATL and SLFNTVATL were both recognized.

HXB2 Location p17 (77-85) Author Location Gag (77-85) Epitope SLYNTVATL Immunogen vaccine

Vector/Type: vaccinia

Species (MHC) human (A2)

References Woodberry et al. 1999

- · A polyepitope vaccine was generated in a vaccinia construct that contiguously encoded seven epitopes, all presented by HLA A-2.
- HHD mice have a transgene of HLA A2 linked to the transmembrane and cytotoxic domains of H-2D<sup>d</sup> – this transgene is the only MHC molecule expressed in the mice.
- CTL responses to Gag (77-85) SLYNTVATL, Pol (476-484) ILKEPVHGV, gp120 (120-128) KLTPLCVTL, and Nef (190-198) AFHHVAREL were observed in HIV polytope HHDvaccinated mice, and these responses were enhanced with vaccinia boost.
- · No CTL immune responses were generated against HLA A2restricted HIV epitopes Nef 157-166 (PLTFGWCYKL), Pol 346-354 (VIYQYMDDL), and Nef 180-189 (VLEWRFD-SRL).

- CTL responses by peptide restimulation in culture with the epitopes selected for inclusion in the polytope - one individual recognized all seven of these epitopes; 7 patients had CTL cultures able to recognize at least one of the epitopes, and 6 of those 7 recognized more than one epitope, but they were not able to test all peptides for all patients; many patients only had three peptides tested.
- SLYNTVATL was recognized by 5/16 HLA-A2 patients.

**HXB2 Location** p17 (77–85) Author Location p17 (77–85) **Epitope** SLYNTVATL Immunogen vaccine

> Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag, gp120, gp41, Protease

Species (MHC) human (A2)

Keywords immunodominance References Carruth et al. 1999

- The vaccine used was a live recombinant canarypox (CP) virus vaccine containing multiple HIV-1 genes (HIV-1 MN gp120, HIV-1 LAI gp41, HIV-1 LAI Gag, HIV-1 LAI protease).
- CD4+ and CD8+ Gag and Env specific CTL responses were detected in only 1/5 vaccinated volunteers, and were not detectable 1 year after vaccination.
- CTL responses to epitopes SLYNTVATL and TVYYGVPVWK from HIV+ control patients were used as positive controls.
- The study explored why vaccinees were non-responsive nonresponse was not due to inherent defects or differences in the ability of these individuals to process and present antigen.
- Lack of response to SLYNTVATL led the authors to speculate that the immunodominance of this epitope in natural infections may not be recapitulated by vaccine antigen.

**HXB2 Location** p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL Epitope name SL9 Immunogen HIV-1 infection Species (MHC) human (A2)

References Birk et al. 1998b

• A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

**HXB2 Location** p17 (77–85) **Author Location** p17 (77–85) **Epitope SLYNTVATL** Epitope name SL9 Immunogen HIV-1 infection Species (MHC) human (A2)

References Callan et al. 1998 • Included as a negative control in a tetramer study of A2-EBV CTL response.

**HXB2 Location** p17 (77–85) **Author Location** p17 **Epitope** SLYNTVATL Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A2)

References Wagner et al. 1998a

• CTL specific for HIV epitopes were used to show that the mediators of both the cytolytic (granzyme A was used as the marker) and non-cytolytic (HIV-1 inhibitory chemokines MIP-1  $\alpha$  and RANTES were used as markers) anti-viral responses are localized within the CTL's cytotoxic granules.

**HXB2 Location** p17 (77–85) Author Location p17 (77–85 HXB2)

**Epitope SLYNTVATL** 

Epitope name SL9 Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2)

References Collins et al. 1998

- Two CTL clones recognize this epitope, but not the NL4-3 form of the epitope SLYNTIAVL.
- Nef down-regulates MHC class I molecules, which inhibits CTL killing, and this down-regulation can be partially compensated for by adding excess soluble peptide.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85)

**Epitope** SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** subtype comparisons

References Durali et al. 1998

- Cross-clade CTL response was studied by determining the CTL activity in seven patients from Bangui, (6 A subtype, and 1 AG recombinant infections) and one A subtype infection from a person living in France originally from Togo, to different antigens expressed in vaccinia.
- Pol reactivity: 8/8 had CTL to A subtype, and 7/8 to B subtype, and HIV-2 Pol was not tested.
- Gag reactivity: 7/8 reacted with A or B subtype gag, 3/8 with HIV-2 Gag.
- Nef reactivity: 7/8 reacted with A subtype, and 5/8 with B subtype, none with HIV-2 Nef.
- Env reactivity: 3/8 reacted with A subtype, 1/8 with B subtype, none with HIV-2 Env.
- Patient B18 had the greatest breadth and diversity of response, and recognized Gag SLYNTVATL and Nef PLTFGWCFKL.

**HXB2 Location** p17 (77–85)

Author Location p17 (77-85)

**Epitope** SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords dendritic cells

References Kundu et al. 1998b

· Allogeneic dendritic cells (DCs) were obtained from HLAidentical siblings, pulsed with rgp160 MN or A2-restricted HIV-1 epitope peptides, and infused monthly into six HIVinfected patients.

- 1/6 showed increased env-specific CTL and increased lymphoproliferative responses, 2/6 showed increase only in proliferative responses, and 3/6 showed no change – pulsed DCs were well tolerated.
- SLYNTVATL is a conserved HLA-A2 epitope included in this study - 3/6 patients had this sequence as their HIV direct sequence, one had the form SLYNTVAVL and all four of these had a detectable CTL response – the other two had either the sequence SLFSAVAVL or SLFSAVAAL and no detectable CTL response.

**HXB2 Location** p17 (77–85)

Author Location p17 (77-85 IIIB)

**Epitope SLYNTVATL** 

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Sipsas et al. 1997

- · HIV IIIB proteins were used to define the range of CTL epitopes recognized by 3 lab workers accidentally infected with HIV-1 IIIB.
- · SLYNTVAVL, a variant found in HIV-1 MANC, was also recognized.
- · SLFNTVAVL, a variant found in HIV-1 NY5CG, was also recognized.

**HXB2 Location** p17 (77–85)

**Author Location** p17

**Epitope** SLYNTVATL

**Epitope name** SL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords subtype comparisons, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades - such crossreactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A subtype consensus is SLfNtvatL.
- The D subtype consensus is SLyNTvATL.

**HXB2 Location** p17 (77–85)

Author Location p17

**Epitope SLYNTVATL** 

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords binding affinity

References Sewell et al. 1997

- · Naturally occurring variants of this epitope escaped killing and acted as antagonists.
- The following variants were found in HIV-1 infected patients who mounted a strong response against this epitope: -F--F--V-, -S---, -SF---, -L---, -I--, -I-V-, -F-I--, -F-I-V-, -F-A---
- All variants bound to A2 with at least half the affinity of SLYNTVATL except the triple mutant: -F-I-V-

• Antagonism could be observed at low concentrations, abrogating lysis at an antagonist: agonist ratio of 1:10 – the antagonism was observed in one SLYNTVATL-specific CTL line but not another.

HXB2 Location p17 (77-85) Author Location p17 (77-85 HXB2) **Epitope SLYNTVATL** Epitope name SL9 Subtype B Immunogen HIV-1 infection Species (MHC) human (A2) Keywords kinetics References Yang et al. 1997b

- · A chimeric universal T cell receptor was created by linking CD4 or an HIV-specific anti-gp41 Ig sequence to the signaling domain of the T cell receptor chain  $\zeta$ , and transduced into CD8+ cells.
- The response using universal-receptor-bearing CD8+ cells to lyse infected cells in vitro was comparable to the natural occurring responses of CTL-clones from HIV+ individuals in terms of kinetics and efficiency.
- · A CTL clone specific for this epitope was used for the compar-

**HXB2 Location** p17 (77–85) Author Location p17 (77-85) **Epitope SLYNTVATL Epitope name** SL9 **Immunogen** in vitro stimulation or selection

Species (MHC) human (A2)

References Stuhler & Schlossman 1997

• Keyhole limpit hemocyanin or tetanus toxoid Th epitope coexpression with peptide CTL epitopes on the same APC was required for induction of peptide-specific CTL.

HXB2 Location p17 (77-85) Author Location p17 (77-85) Epitope SLYNTVATL Epitope name SL9 Immunogen HIV-1 infection Species (MHC) human (A2) References Yang et al. 1996

- CD4+ cell lines acutely infected with HIV were studied to determine their susceptibility to lysis by CTL.
- Clones specific for RT lysed HIV-1 infected cells at lower levels than Env or Gag specific clones.
- The distinction was thought to be due to lower expression of RT relative to Env and Gag.
- CTL can lyse infected cells early after infection, possibly prior to viral production.

**HXB2 Location** p17 (77–85) Author Location p17 (77–85) **Epitope** SLYNTVATL Epitope name SL9 Immunogen HIV-1 infection Species (MHC) human (A2)

Assay type CTL suppression of replication

References Yang et al. 1997a

- comparable to those found in vivo.
- CTL produced HIV-1-suppressive soluble factors MIP-1 $\alpha$ , MIP-1 $\beta$ , RANTES, after antigen-specific activation.
- CTL suppress HIV replication more efficiently in HLAmatched cells.

HXB2 Location p17 (77-85) Author Location p17 (77-85 LAI) **Epitope SLYNTVATL** Epitope name SL9 Subtype B Immunogen HIV-1 infection Species (MHC) human (A2)

References Parker et al. 1992; Parker et al. 1994

· Examined in the context of motifs important for HLA-A2 bind-

**Author Location** p17 (77–85 LAI) **Epitope SLYNTVATL** Epitope name SL9 Subtype B Immunogen HIV-1 infection Species (MHC) human (A2) Keywords review

References McMichael & Walker 1994

• Review of HIV CTL epitopes.

**HXB2 Location** p17 (77–85)

HXB2 Location p17 (77-85) Author Location p17 (77–85) **Epitope** SLYNTVATL Epitope name SL9 Immunogen HIV-1 infection Species (MHC) human (A2)

References Tsomides et al. 1994 • CTL clones recognize naturally processed peptide.

**HXB2 Location** p17 (77–85) **Author Location** p17 (77–85) **Epitope** SLYNTVATL Epitope name SL9

Immunogen in vitro stimulation or selection

Species (MHC) human (A2)

References Stuhler & Schlossman 1997

• A three cell-type cluster consisting of APCs, Th, and CTLs is the minimal regulatory unit required for Th cell-dependent induction of CTLs.

HXB2 Location p17 (77-85) Author Location p17 (77–85) Epitope SLYNTVATL Epitope name SL9 Immunogen HIV-1 infection Species (MHC) human (A2) **Keywords** subtype comparisons References Cao et al. 1997a

• The consensus peptides of B and D clade viruses and some Cs have the sequence SLYNTVATL.

 The consensus peptide of A, and some C strains have SLFNT-VATL, a form that is cross-reactive.

HXB2 Location p17 (77–85) Author Location Gag (77–85) Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection
Species (MHC) human (A2)
References Dyer et al. 1999

- CTL specific responses were measured over a 1.3 to 1.5 year period in members of the Sydney Blood Bank Cohort (SBBC) who had been infected with a natural attenuated strain of HIV-1 which was Nef-defective.
- Some of these patients had prolonged high levels of CTL effector and memory cells despite low viral load.

HXB2 Location p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A2) Keywords escape

References Harrer et al. 1998

- Two overlapping epitopes were recognized in a long-term survivor, restricted by two different HLA molecules, HLA-A11(TLYCVHQR) and -A2 (SLYNTVATL)
- Viral sequence substitutions were present in this individual which did not affect viral replication and did not alter CTLrecognition of the A2 epitope, but reduced recognition of the A11 epitope, indicative of immune escape.

HXB2 Location p17 (77–85)

Author Location p17 (77–85 SF2)

Epitope SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords acute/early infection

References Altfeld et al. 2001a

- The relative contribution of CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- Individuals with long-term nonprogressive and treated chronic HIV-1 infection targeted Vpr more frequently than individuals with treated acute infection.
- Vpr is a frequent target of HIV-1 specific CD8+ T-cells a response was detected in 45% of individuals tested and Vpr and p17 were the most preferentially targeted proteins per unit length by CD8+ T-cells.
- The A2 epitopes Vpr AIIRLLQQL and p17 SLYNTVATL do not account for the dominance of Vpr and p17, the result holds even when HLA-A2+ individuals are excluded.

**HXB2 Location** p17 (77–85) **Author Location** p17 (BRU)

Epitope SLYNTVATL

Epitope name SL9

Immunogen in vitro stimulation or selection

Species (MHC) human (A2)

**Keywords** epitope processing, dendritic cells **References** Buseyne *et al.* 2001

- Exogenous presentation or cross-presentation of epitopes by antigen presenting cells (APC) without protein synthesis is an alternative pathway for CTL epitope processing that may be important in the initial generation of viral specific CTL.
- Dendritic cells treated with AZT to inhibit protein synthesis were able to elicit a strong specific CTL response in SLYNT-VATL specific CTL line EM71-1 without protein synthesis, while macrophages demonstrated a decreased presentation efficiency.
- Exogenous Gag epitope presentation was Env-dependent and required receptor-dependent fusion.

**HXB2 Location** p17 (77–85)

**Author Location** p17

Epitope SLYNTVATL
Immunogen HIV-1 infection
Species (MHC) human (A2)

References Kostense et al. 2001

- HLA tetramers to six epitopes were used to study HLA-A2, B8 and B57 CTL in 54 patients HIV-specific tetramer positive cells were inversely correlated with viral load in patients with high CD4, but in patients with CD4 T-cells below 400 high tetramer frequencies were found despite high viral load.
- Most patients have high levels of HIV-specific T-cell expansions, but many of these cells aren't functional.
- In 15 of the patients, the proportion of IFN gamma producing tetramer cells correlated with AIDS-free survival.
- In one patient with a SLYNTVATL response, no SLYNTVATL mutations were found among 21 clones despite high viral load (260,000 RNA copies/ml serum), suggesting low *in vivo* efficacy of the SLYNTVATL response.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85)

**Epitope** SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p17 (77–85)

Author Location p17

Epitope SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords HAART, ART, immunodominance

References Seth et al. 2001

- CTL responses were studied by tetramer staining in 41 patients with combination therapy activated CD8+ T-cells decline as the viral load drops in response to therapy, but the overall level of antigen-specific cells capable of differentiating into effectors stays constant and new epitopes may be recognized.
- 6/10 A\*0201+ individuals had HIV-specific tetramer staining cells, and 5 of these declined upon successful therapy.
- 4/10 A\*0201+ individuals with chronic HIV-1 infection recognized this epitope.

• Prior to therapy, the mean percentage of CD8+ cells that • Variants SL(F/Y)NTVATL are A/B clade specific. recognized the immunodominant epitope SLYNTVATL was six-fold greater than the percentage recognizing the epitope ILKEPVHGV.

HXB2 Location p17 (77-85) Author Location p17 (77-85) Epitope SLYNTVATL Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A2)

Keywords HAART, ART, TCR usage

References Islam et al. 2001

- Transcript frequencies were followed for four CTL clones from patient 115, with a chronic and stable HIV-1 infection, and tracked in a longitudinal study of samples collected 6-11 years post infection: clone M21 and E15 recognize ERYLKDQQL, and clone D87 recognizes variant ERYLQDQQL, and clone p175b recognizes the A2 epitope SLYNTVATL.
- This epitope sequence from clone p175b uses the V $\beta$ 5, CDR3 (FDS), J $\beta$ 2.7 TCR beta gene.
- Responses were stable even through HAART with undetectable viral loads, but frequencies varied over time by 100-fold, ranging from 0.012% of the total population for SLYNTVATL at its lowest point to 3.78% for M21, with the relative frequencies of clones shifting over time.

HXB2 Location p17 (77-85) **Author Location** p17 (77–85 SF2) **Epitope** SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A2)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- · The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- · Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A2+ individuals that had a CTL response to this epitope broken down by group: 1/10 group 1, 2/6 group 2, and 2/4 group 3.

HXB2 Location p17 (77-85) Author Location p17 (77-85) **Epitope SLFNTVATL** 

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A2)

Keywords subtype comparisons, HIV exposed persistently seronegative (HEPS), immunodomi-

nance

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-A2 women, 1/10 HEPS and 22/26 HIV-1 infected women recognized this epitope, likelihood ratio 18.3, p value < 0.003, and ILK(D/E)PVHGV tended to be more reactive in HEPS women, SL(F/Y)NTVATL in infected women.
- The dominant response to this HLA allele was to this epitope in the 1/10 HEPS case and in 18 of the 22/26 HIV-1 infected women that responded.
- Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.
- Subject ML 1250 had an A2 response to ILK(D/E)PVHGV prior to seroconversion, which switched to SL(F/Y)NTVATL post-seroconversion.
- Subjects ML 1575 and ML 1592 had no response to SL(F/Y)NTVATL prior to seroconversion, but made responses post-seroconversion.
- Subject ML 1760 had an A2 response to ILK(D/E)PVHGV prior to seroconversion, and gained responses to epitopes A2 SL(F/Y)NTVATL and B27 KRWII(L/M)GLNK postseroconversion.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85 93TH253 subtype CRF01)

**Epitope SLYNTIATL** Epitope name G77-85 Subtype CRF01\_AE Immunogen HIV-1 infection Species (MHC) human (A2)

Keywords HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers - weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV positive controls, and 0/9 HIV negative women that were not exposed.
- This epitope was reactive in HIV+ control study subjects 125 and 144 who carried HLA-A2.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85 93TH253 subtype CRF01)

**Epitope SLYNTIATL** Subtype CRF01\_AE **Immunogen** HIV-1 infection **Species** (MHC) human (A2)

 $\textbf{Keywords} \ \ \text{subtype comparisons}$ 

References Bond et al. 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- 2/4 tested FSWs recognized the E clade version of this epitope, SLYNTIATL, the B clade version is SLYNTVATL.
- This epitope was only conserved in CRF01 and subtypes B and D, and exact matches were uncommon.

HXB2 Location p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A2)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- SLYNTVATL was the dominant A2 epitope recognized in patients with chronic infection, except for Subject 11841 who recognized 5/8 epitopes.
- Three subjects had an A2 response only to SLYNTVATL.
- The two subjects with acute infection did not respond to SLYNTVATL.

**HXB2 Location** p17 (77–85) **Author Location** p17 (77–85)

**Epitope** SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A2)

**Keywords** mother-to-infant transmission, escape

References Goulder et al. 2001c

- Immune escape variants in this epitope where transmitted both horizontally and vertically in two families.
- Eight transmitting mothers and 14 non-transmitting mothers were studied and variation within the SL9 epitope was associated carrying HLA-A2 (P=0.04), but no link between variation from the SL9 consensus and vertical transmission was established.

HXB2 Location p17 (77–85) Author Location p17 (SF2)

Epitope SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A2)

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.
- The response to the peptide was CD8 dependent, but the HLA presenting molecule and optimal epitope were not determined.

**HXB2 Location** p17 (77–85)

**Author Location** 

**Epitope** SLYNTVATL

**Epitope name** Gag-SL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA A02, 17/30 (57%) recognized this epitope.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85 LAI)

**Epitope SLYNTVATL** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** HAART, ART, epitope processing, immunodominance

References Kelleher et al. 2001a

- Ritonavir (RTV) inhibits chymotryptic activity in the 20S proteasome *in vitro*, as does Saquinavir (SQV) to a lesser extent; Indinavir (IDV) does not. Thus there is concern protease inhibitors may adversely effect CTL epitope processing, but this paper indicates that processing is not inhibited at therapeutically relevant concentrations of RTV when the proteasome is functioning in an intracellular context.
- RTV did not reduce antigen presentation and concentration of the two immunodominant Gag CTL epitopes (KRWIIMGLNK (B27) and SLYNTVATL (A2)).
- RTV did not inhibit the processing and assembly of HLA-B35 or -A2, which are assembled with a rapid and moderate time course, respectively, or of HLA-A3, -B27 and -B39.

**HXB2 Location** p17 (77–85)

Author Location p17

**Epitope** SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

**HXB2 Location** p17 (77–85)

Author Location p17 (77-85 NL43)

**Epitope** SLYNTVATL

Subtype B

Immunogen HIV-1 infection **Species** (MHC) human (A2)

Keywords class I down-regulation by Nef

References Yang et al. 2002

• Nef down-modulates class I protein expression, and this study demonstrates directly that Nef-deleted HIV-1 NL-43 can be more effectively killed *in vitro* than NL-43 with an intact Nef. The effect was shown to be specific for class I presentation of epitopes, and unlike Nef, deleting Vpr did not alter CTL susceptibility of NL-43. The CTL clone 18030D23, specific for the class I A2 presented SLYNTVATL epitope, was one of four used in this study.

HXB2 Location p17 (77–85)

Author Location p17 (77–85 BRU)

Epitope SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords epitope processing

References Cohen et al. 2002

- The antigen presentation of two A2-restricted epitopes was compared, SLYNTVATL (p17) and ILKEPVHGV (RT). HIV-1 infected cells were more sensitive to lysis by SLYNTVATLspecific CTL than by ILKEPVHGV-specific CTL, because of a higher density of SLYNTVATL-A2 resulting from differences in processing.
- Incubation with a T1-cell proteolytic extract showed that by four hours, 25% of a p17 peptide had a C-term Leu-85 and were SLYNTVATL-precursors, while ILKEPVHGV-precursors were far less frequent (6.8%) even with four times more proteolytic extract after 30 hours.
- p17 was preferentially cleaved between Leu85 and Tyr86, while appropriate Val484 and Tyr485 cleavage was minor for RT.
- In a competition experiment, RSLYNTVATL bound TAP 3.7-fold more efficiently than RT peptides.
- No difference in CTL avidity was detected in six patients with HLA-A2-restricted responses to these epitopes.
- No significant difference in HLA-A2 binding to p17 or RT epitopes was observed.

HXB2 Location p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL

Immunogen vaccine

Strain: B clade IIIB HIV component: Gag, Pol Adjuvant: IL-12

**Species (MHC)** mouse (A2)

References Kmieciak et al. 2001

- Transgenic mice expressing a HLA-A2/Kb chimeric protein were vaccinated with either a p17-p24-p51 fusion protein (vG/P-92) or the Gag-Pol precursor protein (vVK1).
- Compared to vVK1, vG/P-92 induced a significant increase in Gag and Pol induced IFNgamma production and CTL responses, and to the epitopes SLYNTVATL and ILKEPVHGV, as determined by Elispot and 51Cr-release assays.

**HXB2 Location** p17 (77–85)

Author Location Gag (77–85)
Epitope SLYNTVATL
Immunogen HIV-1 infection
Species (MHC) human (A2)
Donor MHC A2, A3, B7, Bw6
Keywords HAART, ART
References Appay et al. 2002

- Four HIV patients with prolonged clinically successful antiviral therapy but with ongoing evidence of replication and Nef mRNA transcription, showed specific T-cell responses by Elispot and Tetramer staining, maintained for 2-4 years after initiation of HAART.
- Nef epitope recognition was detected in all 4 subjects, gp120, Pol and Gag-specific in 1 or 2 subjects.
- The HIV-specific CD8+ T-cells had an intermediate maturation phenotype characterized by low levels of perforin and high levels of CD27 expression.

HXB2 Location p17 (77–85)
Author Location p17 (77–85 NL-43)
Epitope SLYNTVATL
Subtype B
Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** class I down-regulation by Nef, escape **References** Ali *et al.* 2003

- NL43 was passaged in the presence of Nef KEKGGLEGL-specific CTL clones STD11 and KM3, giving rise to rapid selection of escape mutations, including E93G, E93K, K94N+G99R, G95R+G99R, E98K, E98D, G99R, G99E, L100P, and L100I; insertions, deletions, frameshifts and an early stop codon. 34/36 (94%) of sequences carried mutations in the epitope by seven days, 36/36 (100%) by 14 days.
- Several mutations selected by KEKGGLEGL-specific CTL were shown to impair the down regulation of class I MHC by Nef, in particular E93G, E93K, and a truncation mutation at position 51.
- Nef deletion mutants increased 100-fold NL-43 susceptibility to inhibition by CTL specific for epitopes in other proteins, the A2 epitopes ILKEPVHGV in RT and SLYNTVATL in p17 Gag.

**HXB2 Location** p17 (77–85)

**Author Location** Gag

Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Assay type** Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords class I down-regulation by Nef

References Bobbitt et al. 2003

• Nef, through Nef-mediated MHC-1 down-regulation, is not the only viral protein to influence levels of HIV-specific CTL recognition. The Rev L60F mutation, a common natural variant, can decrease CTL recognition of late gene products. The Rev mutation impacts the early to late gene switch, reducing late gene product production (Gag, Pol, Env, Vpu, Vpr and Vif), while increasing Nef production, both of which reduce the impact CTL recognition of late gene products. As expected, Rev L60F rendered HIV infected cells more resistant to CTL that recognized epitopes from the late proteins Env and Gag. Gag expression is reduced more than Env, and Gag-specific CTL were more profoundly affected. Conversely CTL against an epitope in an early gene product, Tat, were more effeciently recognized when infected with viruses carrying the Rev L60F mutation.

 Patients in the asymptomic phase with active immune responses had more CTL resistant viruses, with lower Rev activity, lower Gag expression and greater resistance to Gag-specific CTL killing, while viruses isolated from people with AIDS were more sensitive to CTL killing.

**HXB2 Location** p17 (77–85)

**Author Location** Gag (77–)

Epitope SLYNTVATL

Epitope name Gag77

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Gag Adjuvant: Incomplete Freund's Adjuvant (IFA)

**Species (MHC)** human, transgenic mouse (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This epitope was one of the previously identified HLA-A2 epitopes studied.
- 10/17 HIV-infected HLA-A2+ people in this study recognized this epitope, and CTL and CD8+ T cells responses were elicted by immunization of transgenic mice with this peptide.

**HXB2 Location** p17 (77–85)

**Author Location** p17

Epitope SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A2)

Assay type Intracellular cytokine staining

**Keywords** immunodominance, genital and mucosal immunity

References Kaul et al. 2003

- Predefined immunodominant peptide responses were used to compare CD8+ T-cell responses in the blood and cervix of 16 HIV+ Kenyan sex workers. Cervical responses were detected in 8/10 women from whom adequate samples could be obtained. The frequency of the CD8+ T-cell response in the genital tract was comparable to the blood, with a trend toward being slightly higher.
- The immunodominant response was to this epitope in the PBMC of 1/16 patients (Kaul *et al.* 2001, AIDS, 107:1303).

**HXB2 Location** p17 (77–85) **Author Location** p17

Epitope SLYNTVATL

**Epitope name** SL9

Immunogen HIV-1 infection Species (MHC) human (A2)

**Donor MHC** A2, A24, B38, B60, Cw2, Cw12

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** supervised treatment interruptions (STI), early

treatment

References Montefiori et al. 2003

• HIV-1 + patient AC10 underwent early HAART treatment, which was discontinued 1.5 years later. At this timepoint potent NAb responses against autologous virus were detected. Treatment interruption initially induced weak CD8+ responses directed against 5 epitopes. By days 873d and 923d the CTL response had broadened to target 22 epitopes; of these six were fully characterized. Eventually the virus escaped the NAb response, but escape was not accompanied by a rise in viral load, and the authors suggest the virus was contained by the CTL response.

**HXB2 Location** p17 (77–85)

**Author Location** Gag (77–85)

**Epitope** SLYNTAVTL **Immunogen** HIV-1 infection

Species (MHC) human (A2)

Assay type CD8 T-cell Elispot - IFNγ, Tetramer binding, Flow cytometric T-cell cytokine assay

**Keywords** responses in children **References** Sandberg *et al.* 2003

- 65 vertically HIV-1 infected children, ages 1-16, the majority undergoing ART, were analyzed in regard to their plasma viremia and CD4+ and CD8+ T-cell counts, and CD8+ T-cell responses.
- Using vaccina expressed Gag, Pol, Env, Rev, Nef in target cells in an Elispot assay, 85% of the children recognized at least one HIV antigen. Strong CD8+ T-cell responses were directed against Pol, followed by Gag and Nef. Children younger than 4 had significantly weaker responses (7/14 had no reponse) than older children (only 1/32 had no response, and responses were greater in magnitude).
- SLYNTVATL and ILKEPVHGV tetramers were used to quantitate specific responses. 49 chlidren in an expanded cohort carried HLA-A2. 1/11 children under 3 years of age had detectable CD8+ T-cell responses to SLYNTVATL, 2/11 to ILKEPVHGV. Among children over 3, 11/38 recognized SLYNTVATL and 9/38 recognized ILKEPVHGV.
- Older children that maintained a CD4 count greater than 400 cells/ul tended to have stronger CTL reponses.

**HXB2 Location** p17 (77–85)

Author Location Gag (77–85)

Epitope SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) (A2)

**Donor MHC** A2, A3, B27, B51; A2, A3, B27, B57; A2,

A23, B57

Assay type Cytokine production, CD8 T-cell Elispot -

IFN $\gamma$ , Tetramer binding, Intracellular cyto-

kine staining

**Keywords** assay standardization/improvement, memory cells

References Sun et al. 2003

- This study compares assay methods for testing CTL responses using samples from 20 HIV+ patients. The study compares ELISpot, tetramer-binding, and intracellular IFNγ. Tetramer-binding analysis was performed with Gag (SLYNTVATL) or Pol (ILKEPVHGV) tetramers. Antigen presentation using recombinant vaccinia viruses (rVVs) encoding HIV-LAI Gag, Pol, Env, Nef, Tat and Vif proteins was compared to peptide panels. HIV antigen recognition in memory CTLs was measured by chromium release assay and compared to effector/memory CD8+ T-cells in an IFN-γ ELISpot assay.
- Results: IFNγ Elispot and flow cytometry gave similar frequencies of HIV specific CD8+ T-cells. Tetramer-binding analysis was most sensitive. Pools of peptides and the sum of frequencies of individual peptides were comparable. Elispot assays using peptides were more sensitive than assays using vaccinia expressed proteins. Cr release and Elispot against rVVs gave comparable memory cell responses 2/3s of the time.
- 3/7 HLA-A2+ patients recognized this epitope.

HXB2 Location p17 (77–85) Author Location p17 (77–85 NL43)

Epitope SLYNTVATL

**Epitope name** SL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Assay type** Chromium-release assay, CTL suppression of replication

**Keywords** escape, TCR usage **References** Yang *et al.* 2003a

- Virus was cultured in the presence of CTL lines specific for 5 different epitopes (SLYNTVATL, ILKEPVHGV, IEIKDTKEAL, SEGATPQDL, and KEKGGLEGL) to study the emergence of escape mutations. Escape varied between clones for the same epitope, and between different epitopes. Gag and RT epitope escape, if it occurred at all, tended to be monoclonal and within the epitope, indicating strong fitness constraints, while the Nef epitope escape was rapid, polyconal, and sometimes the result of upstream frameshifts.
- Three CTL clones were studied that recognized SLYNTVATL, 161JxA14, 18030D23, and 115DEC4. The different TCR usage on the CTL clones resulted in different patterns of recognition and escape. 161JxA14 suppressed the variant slFntvatl, 18030D23 did not; conversely the variants slfntIaVl and slFntIatl were suppressed by 18030D23, but not 161JxA14.
- After two weeks of passage the predominant escape mutant from 161JxA14 was slyntIatl. Amino acid residues flanking SL9 were unchanged. Escape mutations did not occur within two weeks for the two additional SL9-specific CTL clones 18030D23 and 115DEC4.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (43)

Epitope SLYNTVATL

Epitope name SL9

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (A2)

Assay type CTL suppression of replication

Keywords class I down-regulation by Nef, early-

expressed proteins, kinetics

References Ali et al. 2004

 Translocation of the gag SLYNTVATL epitope into the early expressed Nef protein resulted in increased antiviral efficiency of SL9 specific CTLs in culture and the loss of MHC-I downregulation by Nef, indicating that both the timing of epitope expression and reduction of MHC-I affect the ability of CTLs to supress HIV-1.

**HXB2 Location** p17 (77–85)

Author Location Gag (77–85 B con)

Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords variant cross-recognition or cross-

neutralization

References Draenert et al. 2004c

- CD8+ T-cell responses that persisted in individuals with advanced disease were studied; most of the optimal epitopes defined were recognized with intermediate to high avidity. On average 13 (range, 2-39) epitopic regions were targeted in an average of 6 proteins (range, 1-8). HAART resulted in decrease in antigen and reduction in gamma IFN EliSpot responses, suggesting active responses to autologous virus. The lack of mutations within most viral epitopes suggest that persistent CTL through late disease do not exert strong immune selection pressure, yet the Elispot assays show robust responses, suggesting to the authors that gamma IFN-based screening methods may not reveal functional CD8+ T-cell impairment in patients with AIDS.
- 3 subjects recognized this epitope with high functional avidity.
   Relative to consensus, 2 individuals that had the SLYNTVATL epitope carried a R -> K mutation proximal to but outside the epitope; possible processing implications were not studied here

**HXB2 Location** p17 (77–85)

**Author Location** Gag

**Epitope** SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

Keywords HIV exposed persistently seronegative

(HEPS)

References Koning et al. 2004

 A high-risk seronegative group of 29 patients showed reduced in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection. • 2/11 HLA A2+ infection-resistant men, compared to 7/9 men pre-seroconversion who went on to become infected, reacted to this epitope.

HXB2 Location p17 (77-85) Author Location p17 (77–85) **Epitope** SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A2) Country Spain.

> Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 14/19 patients recognized this epitope, it was the most commonly recognized of 9 HLA A\*02 epitopes tested.

**HXB2 Location** p17 (77–85) Author Location p17 (77-85)

**Epitope SLYNTVATL** 

Epitope name SL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type Chromium-release assay

Keywords binding affinity, TCR usage, characterizing CD8+ T cells

References Yang et al. 2003b

- Killing efficiency of CTLs and their ability to suppress viral replication is shown to depend more on epitope specificity than to antigenic avidity. Different clones recognizing the same epitope had similar killing efficiency despite their variation in avidity. Nef specific CTL clones tended to be most inhibitory, followed by Gag, then by RT specific clones, regardless of avidity.
- 3/14 CTL T-cell clones tested were specific for Gag/p17-SL9. Under conditions of excess peptide (100ug/ml), there was no difference in their lytic potential; all possessed similar effector capacity. Avidity was measured as the sensitizing dose of peptide required for 50% of maximal killing (SD50), which varied from 20 pg/ml to 100 ng/ml, over four orders of magnitude for all 14 epitopes. The SD50 range for Gag/p17-SL9 was 1,000 -20,000 pg/ml.

HXB2 Location p17 (77-85) **Author Location** Gag (77–85) **Epitope** SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A2)

Assav type Cytokine production, proliferation, CD8 Tcell Elispot - IFNy, Tetramer binding, CD4 T-cell Elispot - IFNγ, Intracellular cytokine staining, Flow cytometric T-cell cytokine as-

Keywords HAART, ART, memory cells, characterizing CD8+ T cells

References Daniel et al. 2004

• CD4+ and CD8+ responses in chronically HIV-1 infected patients on HAART were weak with decreased polyclonality. Only 33% of patients had CD4+ T-cells that could proliferate, and only 22% had HIV-specific CD8+ T-cell responses, and those rare responses showed low perforin levels and persistent expression of CD27, indicating incomplete differentiation and loss of lytic function.

**HXB2 Location** p17 (77–85) **Author Location** p17 **Epitope SLYNTVATL** 

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A2)

Country United States.

Assay type proliferation, Tetramer binding, T-cell Elispot Keywords acute/early infection, characterizing CD8+ T cells, immune dysfunction

References Lichterfeld et al. 2004a

- HIV-1 specific CD8+ T-cells in acute and long-term nonprogressive HIV-1 infection show strong ex-vivo proliferative capacities which are rapidly lost in chronic HIV-1 infection. The loss of CD8+ T-cell function is closely linked with the loss of HIV-1 specific, IL2 secreting CD4+ T-cells. The function can be rescued in vitro and in vivo by restoring the specific CD4+ T-cell help.
- Despite being detectable at high frequencies, CD8+ T-cells specific for SL9 epitope were shown to entirely lose their proliferative capacity in chronic HIV-1 infection. This activity could be restored by co-stimulation with CD4+ T cells isolated from acute infection in an IL-2 dependent manner.

**HXB2 Location** p17 (77–85) **Author Location** Gag (77–85)

**Epitope** SLYNTVATL

**Epitope name** gag 77-85

Subtype B

Immunogen HIV-1 infection, HIV-2 infection

Species (MHC) human (A2)

Country Gambia.

Assay type Tetramer binding, Intracellular cytokine stain-

Keywords escape, TCR usage, variant cross-recognition or cross-neutralization, characterizing CD8+ T cells

References Lopes et al. 2003

• CD8+T cells from HIV-2 infected patients had more polyclonal TCR responses than HIV-1 infected patients, who tended to have oligoclonal responses. This results in limited plasticity of T cell responses to amino acid substitutions within epitopes in HIV-1 infections. HIV-2-specific CD8+ T-cells showed a more diverse TCR usage associated with enhanced CD8 expansion and IFN-gamma production on cross-recognition of variant epitopes.

• Responses to this epitope were characterized in detail. One patient's response to SL9 A2-SLYNTVATL tetramers was shown to have only Vbeta5.1 clonotypes. The naturally occuring HIV-2 variant: slFntvCVI, was not recognized well by this response or by the SLYNTVATL reactive CD8+ T cells in four additional A2+ HIV infected asymptomatic individuals. The subtype A variant, slFntvatl was also poorly recognized, and 4/5 Ala substitutions abrogated responses. All variants bound to HLA-A2 with higher affinity than the index peptide except slyntAatl, which was slightly reduced, so the lack of cross-reactivity must have been due to the TCR.

HXB2 Location p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A2)

Assay type Chromium-release assay

Keywords assay standardization/improvement

References Lubong et al. 2004

 Using IL7 or IL15 in culturing of HIV-1-specific CTL clones was inferior to using IL-2 alone; the addition of these cytokines to IL-2 did not show any advantage. Neither proliferation, survival, nor lytic capacity of HIV-1-specific CTLs was significantly enhanced by addition of IL7 or IL15.

HXB2 Location p17 (77-85)

Author Location p17

Epitope SLYNTVATL

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2)

**Donor MHC** A\*02, A\*30, B\*4402, B\*15

Assay type Tetramer binding, T-cell Elispot

**Keywords** HIV exposed persistently seronegative (HEPS)

References Missale et al. 2004

- HIV-specific T-cell response was tested in patients exposed to blood from a patient with highly replicating HIV; these patients were nosocomially infected with HBV, but uninfected with HIV. HIV-specific T-cell responses were directed to structural and non-structural HIV proteins in 2 patients, suggesting that the virus replicated in these patients sufficiently to prime a cell-mediated immune response that protected them from HIV infection.
- This patient responded to 4/8 HIV epitopes tested in an IFNgamma EliSpot assay or tetramer assay. Responses were detected 8 and 28 weeks after exposure. No response to SLYNT-VATL was detected by either assay.

**HXB2 Location** p17 (77–85)

Author Location p17

**Epitope SLYNTVATL** 

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2)

Country United Kingdom.

**Assay type** Tetramer binding, T-cell Elispot, Intracellular cytokine staining

**Keywords** rate of progression, acute/early infection, characterizing CD8+ T cells, immune dysfunction

References Papagno et al. 2004

Acute HIV-1 infection induces massive activation of HIV-specific and non-HIV-specific CD8+ T-cells resulting in differentiation of these cells. High differentiation of CD8+ T-cells is correlated with disease progression. Differentiation is a natural process but it can be driven by elevated immune activation, such as in HIV infection.

HXB2 Location p17 (77–85)
Author Location p17 (77–85)
Epitope SLFNTVATL
Epitope name SLF
Immunogen HIV-1 infection

**Species (MHC)** human (A2) **Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401, Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

DO6

Assay type CD8 T-cell Elispot - IFNγ, Tetramer binding, CD4 T-cell Elispot - IFNγ

**Keywords** rate of progression, immunodominance, escape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant eptiopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This epitope was one of six epitopes found to be under positive selection for escape mutations and was completely replaced by escape variants between days 327 and 635 (slYntvatl and slYnAvatl).

**HXB2 Location** p17 (77–85)

**Author Location** Gag

**Epitope** SLYNTVATL

**Epitope name** SL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.

• Two escape mutations, at positions 3 and 8, slFntvatl and slynt-vaVl, were found in the most polymorphic residues in the epitope. These were shared between clades B and C. One escape mutation, at position 6, slyntIatl, was found not to correspond to the most polymorphic residue in the epitope.

HXB2 Location p17 (77–85)
Author Location p17 (77–85)
Epitope SLYNTVATL
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (A2)
Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

• The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This is the most commonly recognized A2 epitope in chronic infection, recognized in 62% of 74 A2+ people, but it is rarely recognized in acute infection (in only 1/14 cases).

HXB2 Location p17 (77–85)

Author Location p17 (77–85 HXB2)

Epitope SLYNTVATL

Epitope name 17A

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: multiple epitope immunogen HIV component: p17/p24 Gag,

Pol Adjuvant: IL-12

**Species (MHC)** transgenic mouse (A2)

Assay type Cytokine production, CD8 T-cell Elispot -  $IFN\gamma$ , Chromium-release assay

**Keywords** vaccine-specific epitope characteristics, vaccine antigen design

References Bolesta et al. 2005

- Immunization of transgenic mice with a codon-optimized hGagp17p24-Polp51 DNA plasmid, consisting of clusters of highly conserved CTL epitopes presented by multiple MHC class I alleles, induced 2- to 5-fold higher CD8+ T-cell responses than the corresponding full-length proteins. The modified proteins had the ribosomal frameshift deleted, as well as the potentially immunosuppressive p15, and protease and integrase. This correlated with higher protection against challenge with Gag and Pol expressing recombinant vaccinia virus. Mice immunized with the hGagp17p24-Polp51 also showed an elevated level of type 1 cytokine production as well as an increased titer of p24- and RT-specific IgG2 antibody responses.
- This was 1 of 4 A2 gag/pol epitopes tested. Transgenic mice immunized with the deleted construct induced more potent EliSpot reactions to this epitope than those immunized with full length Gag/Pol.

HXB2 Location p17 (77–85) Author Location p17 (77–85) Epitope SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A2)

Donor MHC A2, A2, B44, B70; A2, A31, B51, B58w4

Country United States.

**Assay type** Intracellular cytokine staining, Flow cytomet-

ric T-cell cytokine assay

Keywords HAART, ART, escape, variant cross-

recognition or cross-neutralization

References Casazza et al. 2005

- Evidence of continued viral evolution during HAART therapy despite low viral load was found in 1/5 patients studied.
- This epitope was recognized in 2 individuals and was invariant in both prior to HAART (20/20 clones in each).

**HXB2 Location** p17 (77–85)

**Author Location** 

Epitope SLYNTVATL Immunogen HIV-1 infection Species (MHC) human (A2)

Country Germany.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, escape, variant cross-recognition or cross-neutralization, optimal

epitope

References Harrer et al. 2005

- An HLA-B13-restricted optimal epitope was defined in Nef, R19. The frequency of CTLs specific for this epitope in B13positive patients exceeded the number of CTLs against other epitopes, indicating that this is a dominant epitope in B13positive subjects. Three B13-positive patients who had an immunodominant response to this epitope were good controllers of their infection, with low viral loads over long periods.
- Five A2+ B13+ patients were found to make an immunodominant response to the B13 epitope RI9. 0/5 recognized ILKEPVHGV, and only 1/5 recognized SLYNTAVTL, with a much lower frequency than the B13 response.

**HXB2 Location** p17 (77–85)

Author Location Gag (77–85 BRU)

Epitope SLYNTVATL

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 3/9 CRF02\_AG-infected patients, and by 2/9 B-infected patients.

**HXB2 Location** p17 (77–85)

**Author Location** Gag

Epitope SLYNTVATL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Netherlands.

Assay type Tetramer binding, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, rate of progression, escape, characterizing CD8+ T cells

References Jansen et al. 2005

- HLA-B57 has been associated with long term non-progression in HIV+ people. The number and responsiveness of CD8 T-cells directed to different Gag peptides presented by HLA-A2, -B8 and B57 were compared. T cells specific for the HLA-B57 epitope KAFSPEVIPMF responded to a higher extent and more readily to antigenic stimulation than those specific for the A2 epitope SLYNTVATL and the B8 epitope EIYKRWII.
- In 3/4 A2 subjects that were sequenced, epitope variants dominated: 2 subjects carried slFntvatl, and the other slyntIatl.
- Tetramer decay experiments indicate that the HLA-B57 peptide has a higher half-life than the A2 and B8 peptides. The authors point out that CD8+ T cells with high binding affinity may require less help.

**HXB2 Location** p17 (77–85) **Author Location** Gag (77–85)

Epitope SLYNTVATL

Epitope name SL9 Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2)

**Donor MHC** A\*02, A\*68, B\*14, B\*52, Cw\*08, Cw\*12

Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** escape, optimal epitope **References** Koibuchi *et al.* 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- Elispot responses to the consensus form of this epitope, SLYNT-VATL, were much more intense than to the most common variants of the epitope found over time in this individual, SLfN-TiATL and SLfNTVAvL; these may be escape variants. The strong response to the consensus form persisted, despite the fact it was not observed among the autologous sequences during 6 years of chronic infection.

**HXB2 Location** p17 (77–85)

Author Location p17 (77–85)

**Epitope SLYNTVATL** 

Epitope name SL9

Subtype B

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A2)

Country Canada.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance, genital and mucosal immunity, characterizing CD8+ T cells

References Makedonas et al. 2005

- CD8 T-cell responses were studied in individuals who remained seronegative in spite of being mucosally (group 1) or intravenously (group 2) exposed to HIV-1. A similar proportion of subjects from each group recognized at least 1 HIV peptide, and they recognized peptides with similar cumulative intensity. The proportion of responding individuals in both groups was significantly greater than in a low-risk, negative control group.
   One exposed uninfected subject recognized 7 epitopes.
- HLA-A\*0201 epitopes that are immunodominant in chronically infected individuals were rarely stimulatory in exposed uninfected individuals. SLYNTVATL was recognized by one HLA A2+ individual in each group (1/11 vs 1/5), while none of the exposed uninfected individuals tested responded to ILKEPVHGV. In contrast, chronically infected subjects recognized these epitopes at a frequency of 69% and 31%, respectively.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85)

Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Germany.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

**Keywords** HAART, ART, characterizing CD8+ T cells,

optimal epitope

References Schmitt-Haendle et al. 2005

- CTL responses to 3 HLA-A2-restricted epitopes were investigated in 51 HIV-1 infected HLA-A2+ individuals. The most prevalent response was seen for IV9, followed by SL9. The VL9 epitope was not recognized. There was a significant correlation of CTL activity to the CD8 counts in peripheral blood, but no correlation to CD4 counts, viral load, or antiviral therapy.
- SL9 was only recognized in 13.7% of the individuals tested.

**HXB2 Location** p17 (77–85)

**Author Location** p17

Epitope SLYNTVATL

Epitope name SL9

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2, A\*0202)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found B clade epitopes are often cross-reactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among B and D clade viruses.

- The Clade A version of the epitope, SLFNTVATL, was preferentially recognized by CTL.
   Longitudinal study of 8 infants with prolonged viral suppression due to combination antiretroviral therapy showed no HIV-1
- This epitope was recognized by two different exposed seronegative prostitutes.

**HXB2 Location** p17 (77–85)

Author Location p17

**Epitope** SLYNTVATL **Immunogen** HIV-1 infection

Species (MHC) human

**Keywords** review, escape

References Sewell et al. 2000

 Review of the impact of CTL on viral immunity and escape that notes that SLYNTVATL-tetramer binding cells in individuals that react to this epitope inversely correlate with plasma viral load.

**HXB2 Location** p17 (77–85)

Author Location (SF2, HXBc2/Bal chimeric)

**Epitope** SLYNTVATL

Epitope name SL9

Subtype B

Immunogen HIV-1 infection

Species (MHC)

**Keywords** rate of progression, escape

References Douek et al. 2002

- Seven HIV-positive subjects tended to make their strongest CD8+ T-cell response against Gag; these responses had varying breadth and magnitude that were unrelated to disease progression
- Patient TX7 primarily recognized SL9 during a three year study period and used six T-cell clonotypes for this recognition.
- SLYNTVATL was the only form of the epitope found initially, but three alternate forms eventually appeared: SLYNTVAVL, SLYNTIATL, and most commonly SLYNTIAVL. These distinct forms bind A2, but have distinct abilities to stimulate different T-cell clonotypes.
- In subject TX7, the observed mutations of SL9 failed to escape overall CTL recognition, presumably because the six T-cell clonotypes allowed a more flexible response.
- The BV17 T-cell clone recognized SL9 but not SLYNTIAVL, and BV17 became undetectable at week 20 when SLYNTI-AVL predominated. Subsequently BV17 became the second most common clone. Thus the relative frequency of the T-cell clonotypes varied with respect to each other and to epitope variation.

**HXB2 Location** p17 (77–85)

**Author Location** p17 (77–85 LAI)

**Epitope** SLYNTVATL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A\*0201

Keywords HAART, ART, responses in children

References Luzuriaga et al. 2000

- Longitudinal study of 8 infants with prolonged viral suppression due to combination antiretroviral therapy showed no HIV-1 specific CTL responses in peripheral blood cells. 6/8 were studied using a Chromium release assay and no response was detected using Gag expressed in vaccinia in the target cells. Three HLA-A\*0201 children were tested using SLYNTVATL or ILKEPVHGV HLA A\*0201 tetramers and again no HIV-specific response was detected, either using PBMC specimens, or PBMC which had been stimulated *in vitro* for a week.
- In contrast, one of the children with therapy suppressed HIV viral replication who was co-infected with HIV and EBV, while HIV-tetramer negative, had EBV-tetramer staining cells at a frequency of 0.14% in the PBMC.

HXB2 Location p17 (77-86)

**Author Location** Gag

Epitope SLYNTVATLY

Epitope name 1261

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A01, A02, B08, Cw16; A02, A30, B35, B49,

Cw04, Cw07; A02, A03, B7, B58, Cw07;

A02, A03, B08, B51, Cw01, Cw07 **Country** United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for SLYNTVATLY: 78%

**HXB2 Location** p17 (77–91)

**Author Location** p17 (77–85)

Epitope SLYNTVATLYCVHQR

Subtype A, D

Immunogen HIV-1 infection

**Species (MHC)** human (A\*3002, A\*0201)

**Donor MHC** A\*3002, A\*6801, B\*5703, B\*5802; A\*0201,

A\*2902, B\*1402, B\*1503

Country Uganda.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, variant cross-

recognition or cross-neutralization

References Barugahare et al. 2005

 T-cell immune responses were examined in HIV-1 infected Ugandans. Similar levels of cross-clade responses for Gag, Env and Nef were observed. Higher frequencies of responses in conserved regions were found for Gag within a single clade, while areas with higher viral sequence variation had fewer detectable responses. Differential interclade sequence homologies between the Gag regions did not affect the respective level of cross-clade recognition.  The sequence contains a known A2 epitope and a known A\*3002 epitope, and the subjects recognizing it each carry an HLA with a previously-defined restriction. The viral sequence isolated from the subjects was slFntvatlycvhqr, and was reactive.

**HXB2 Location** p17 (78–85)

Author Location p17

**Epitope** LYNTVATL

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (A24)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on a patient with an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

**HXB2 Location** p17 (78–85)

**Author Location** 

Epitope LYNTVATL

Immunogen

Species (MHC) human (Cw\*14)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an Cw14 epitope.

**HXB2 Location** p17 (78–86)

Author Location (C consensus)

 ${\bf Epitope} \ \ {\tt LYNTVATLY}$ 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*29)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the Y2 residue of LYNTVATLY are associated with the presence of the HLA presenting molecule in the host.

HXB2 Location p17 (78-86)

Author Location p17 (78–86)

**Epitope** LYNTVATLY

Immunogen

Species (MHC) human (A\*2902)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an A\*2902 epitope.

**HXB2 Location** p17 (78–86)

**Author Location** (78–86)

**Epitope** LFNTVATLY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2902)

Assay type Other

Keywords HLA associated polymorphism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- LFNTVATLY was a previously defined A\*2902 presented epitope that encompassed an A\*29 associated polymorphism, LfNTVATLY,in the second position.

**HXB2 Location** p17 (78–86)

Author Location p17

Epitope LYNTVATLY

Epitope name LY-9

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (A\*2902, B\*4403)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Eli-

spot - IFNγ, Intracellular cytokine staining,

Chromium-release assay

**Keywords** subtype comparisons, epitope processing, immunodominance, cross-presentation by differ-

ent HLA

References Masemola et al. 2004b

 Highly targeted regions in Gag for CD8+ T-cells were defined for individuals with C clade infections in South Africa. 9 specific epitopes within the most reactive regions were characterized. This is 1 of 5 novel epitopes that were found among subtype C HIV-1 from African patients who hadn't previously been identified in B clade infections. Some epitopes were shown to be promiscuous, presented by multiple class I restricting alleles. • LYNTVATLY was presented by A\*2902 and B\*4403. B\*44 is more common among Caucasians than Zulus (allele frequency 0.149 versus 0.107), while A\*29 is more common in Zulus (0.045 versus 0.125).

HXB2 Location p17 (78–86)
Author Location (C consensus)
Epitope LYNTVATLY
Subtype C

Immunogen HIV-1 infection Species (MHC) human (A29) Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$  Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p17 (78–86)

**Author Location** 

**Epitope** LYNTVATLY

Immunogen

Species (MHC) human (B\*4403)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes that this is an B\*4403 epitope.

HXB2 Location p17 (80–88) Author Location Gag (80–) Epitope NTVATLYCV

Epitope name Gag80

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: p17 Gag Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human, transgenic mouse (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was an intermediate A2 binder, and induced CTL and CD8+ T-cell IFN gamma responses in mice. Responses were detected in 1/17 HIV+ HLA-A2 subjects.

HXB2 Location p17 (82-91)

**Author Location** p17 (82–91 93TH253 subtype CRF01)

Epitope IATLWCVHQR Epitope name G82-91 Subtype CRF01\_AE

Immunogen HIV-1 infection, HIV-1 exposed seronegative

**Species (MHC)** human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was weakly reactive in the HEPS study subject 265 who was HLA A2/A11.
- This epitope was strongly reactive in HIV+ study subject 053 who carried HLA-A11.

HXB2 Location p17 (82-91)

Author Location p17 (82–91 93TH253 subtype CRF01)

Epitope IATLWCVHQR
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A11)

**Keywords** subtype comparisons

References Bond et al. 2001

- HLA-A11 CRF01 (called subtype E in Bond *et al.*) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This epitope was predicted by the EpiMatrix method to be likely to bind to A11, and it served as an epitope in the FSWs, it was one of the six A11 epitopes that had been previously defined.
- 3/8 tested FSWs recognized this epitope.
- This epitope was not conserved in other subtypes, and exact matches were uncommon.

**HXB2 Location** p17 (84–91)

**Author Location** Gag (83–90)

Epitope TLYCVHQR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*1101)

Keywords subtype comparisons, TCR usage

References Fukada et al. 2002

- Counterparts for eight known clade B HLA A\*1101 epitopes were generated for clade E (CRF01). Three epitopes, identical among clade A-E, were cross-reactive and recognized by clade E infected individuals. The clade E and B analogs to three more HLA A\*1101 epitopes was recognized in a clade specific manner. Two other HLA A\*1101 clade B defined epitopes were found not to have stimulated a response in clade E infected individuals.
- TLYCVHQR was found to elicit clade-specific responses in clade B (TLYCVHQR is most common, and is also common in clade A the variant tlycvhqK is common in clade B) and clade E (tlWcvhqr is most common). TLYCVHQR was not recognized by any CTL, tlycvhqK was recognized by CTL from 1/5 B clade infected Japanese subjects, and tlWcvhqr was not recognized by CTL from infected Thai subjects, so this seems to be a B clade exclusive epitope.
- The binding of the variant peptides to HLA A\*1101 was comparable, but CTL that recognized tlycvhqK did not crossrecognize the other forms, implicating TCR interaction differences.

HXB2 Location p17 (84–91)
Author Location p17 (83–91)
Epitope TLYCVHQR
Immunogen HIV-1 infection
Species (MHC) human (A11)
Keywords escape
References Harrer et al. 1998

- Two overlapping epitopes were recognized in a long-term survivor, restricted by two different HLA molecules, HLA-A11(TLYCVHQR) and HLA-A2 (SLYNTVATL)
- Viral sequence substitutions were present in this individual which did not affect viral replication and did not alter CTLrecognition of the A2 epitope, but reduced recognition of the A11 epitope, indicative of immune escape.
- A Q90E substitution resulted in a loss of the ability of the peptide to induce lysis, a R91K substitution was still reactive, and a R91Q substitution showed a reduced ability to stimulate lysis.

HXB2 Location p17 (84–92)
Author Location p17 (84–92)
Epitope TLYCVHQRI
Immunogen HIV-1 infection
Species (MHC) human (A\*1101)
Keywords optimal epitope
References Frahm et al. 2007

• C. Brander notes that this is an A\*1101 epitope.

HXB2 Location p17 (84–92)
Author Location Gag (83–91 SUMA)
Epitope TLYCVHQKI
Epitope name Gag TI9
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (A\*1103)
Donor MHC A\*1103, A\*2402, B\*1402, B\*1501, Cw\*0802
Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** dynamics, acute/early infection, characterizing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4
  T-cell counts through seven years of follow up. In contrast to
  more rapid progressors, WEAU and BORI, SUMA a broad
  response to 24 epitopes, with little immunodominance. Two
  peptides were somewhat more intensely recognized in acute
  infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

HXB2 Location p17 (84–92) Author Location p17 (84–92) Epitope TLYCVHQRI

Immunogen HIV-1 infection Species (MHC) human (A11)

**Keywords** responses in children, mother-to-infant transmission

References Brander & Walker 1995

• Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.

HXB2 Location p17 (84–92) Author Location p17 (84–92) Epitope TLYCVHQRI Immunogen HIV-1 infection Species (MHC) human (A11) References Birk et al. 1998b

 A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

HXB2 Location p17 (84–92)
Author Location p17 (84–92)
Epitope TLYCVHQRI
Immunogen HIV-1 infection
Species (MHC) human (A11)
References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location p17 (84–92) Author Location p17 (84–92 SF2) Epitope TLYCVHQRI **Immunogen** HIV-1 infection **Species** (MHC) human (A11)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A11+ individuals that had a CTL response to this epitope broken down by group: 0/3 group 1, 0/0 group 2, and 1/2 group 3.

**HXB2 Location** p17 (84–92)

Author Location p17 (84-92)

Epitope TLYCVHQRI

Immunogen HIV-1 infection, HIV-1 exposed seronegative

 $\textbf{Species} \ (\textbf{MHC}) \ \ \text{human} \ (A11)$ 

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** p17 (85–92)

**Author Location** p17

Epitope LYCVHQKI

Subtype D

**Immunogen** HIV-1 infection **Species (MHC)** human (A24)

Decies (MHC) human (A24)

Donor MHC A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on a patient with an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

HXB2 Location p17 (86-101)

Author Location p17 (SF2)

Epitope YCVHQRIEIKDTKEAL

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.
- The response to the peptide was CD8 dependent, but the HLA
  presenting molecule and optimal epitope were not determined.

**HXB2 Location** p17 (86–101)

Author Location p17 (SF2)

Epitope YCVHQRIEIKDTKEAL

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.
- The response to the peptide was CD8 dependent, but the HLA presenting molecule and optimal epitope were not determined.

**HXB2 Location** p17 (87–105)

Author Location p17 (91–105 SF2)

Epitope CRIDVKDTKEALEKIE

Immunogen HIV-1 infection

Species (MHC) human

**References** Lieberman *et al.* 1997b

 CTL expanded ex vivo were later infused into HIV-1 infected patients.

HXB2 Location p17 (88–115)

**Author Location** p17 (88–115 ARV)

Epitope VHQRIEIKDTKEALDKIEEEQNKSKKKA

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Achour et al. 1990

• B cell epitope HGP-30 also serves as a CTL epitope.

**HXB2 Location** p17 (88–115)

Author Location p17 (88-115 ARV)

Epitope VHQRIEIKDTKEALDKIEEEQNKSKKKA

Immunogen vaccine

Vector/Type: peptide HIV component: CD4BS, HPG30, V3 Adjuvant: IL-12

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hamajima et al. 1997

- B cell epitope HGP-30 also serves as a CTL epitope.
- Vaccine combined HGP-30, V3 loop peptide variants, and CD4 binding site peptide.
- IL-12 expression plasmid included with the vaccination enhanced the CTL response.

**HXB2 Location** p17 (91–101)

**Author Location** p17 (SF2)

Epitope RIDVKDTKEAL

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons, immunodominance **References** Goulder *et al.* 2000a

- The CTL-dominant response was focused on this epitope in an HIV+ Haitian living in Boston, who was A23/68 B45/72 Cw2/16 – this epitope fell outside the most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

HXB2 Location p17 (91–105) Author Location p17 (91–105 SF2) Epitope RIDVKDTKEALEKIE Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A3, A24, B8, B55.

HXB2 Location p17 (92–101)
Author Location p17 (92–101)
Epitope IEIKDTKEAL
Immunogen HIV-1 infection
Species (MHC) human (B\*4001)
Keywords optimal epitope
References Frahm et al. 2007

• C. Brander notes this is a B\*4001 epitope.

HXB2 Location p17 (92-101)

Author Location p17

Epitope IEIKDTKEAL
Immunogen HIV-1 infection
Species (MHC) human (B60)

**References** Wagner *et al.* 1998a

 CTL specific for HIV epitopes were used to show that the mediators of both the cytolytic (granzyme A was used as the marker) and non-cytolytic (HIV-1 inhibitory chemokines MIP-1 α and RANTES were used as markers) anti-viral responses are localized within the CTL's cytotoxic granules.

HXB2 Location p17 (92–101) Author Location p17 (92–101 SF2) Epitope IEIKDTKEAL Immunogen HIV-1 infection Species (MHC) human (B60)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B60+ individuals that had a CTL response to this epitope broken down by group: 0/2 group 1, 1/1 group 2, and 0/0 group 3.

HXB2 Location p17 (92–101)
Author Location p17 (SF2)
Epitope IEIKDTKEAL
Immunogen HIV-1 infection
Species (MHC) human (B60)
References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes.
- B60 is present in 10-20% of the Caucasoid and very common in Asian populations.

**HXB2 Location** p17 (92–101)

Author Location Gag (92-101)

Epitope IEIKDTKEAL

Immunogen HIV-1 infection

Species (MHC) human (B60)

Keywords class I down-regulation by Nef

References Yang et al. 2002

 Nef down-modulates class I protein expression, and this study demonstrates directly that Nef-deleted HIV-1 NL-43 can be more effectively killed *in vitro* than NL-43 with an intact Nef. The effect was shown to be specific for class I presentation of epitopes, and unlike Nef, deleting Vpr did not alter CTL susceptibility of NL43 infected cells. The CTL clone 161JD27, specific for the class I B60 presented epitope IEIKDTKEAL, was one of four used in this study.

**HXB2 Location** p17 (92–101)

Author Location p17 (92-101 NL43)

Epitope IEIKDTKEAL

Epitope name IL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B60)

**Assay type** Chromium-release assay, CTL suppression of replication

Keywords escape

References Yang et al. 2003a

Virus was cultured in the presence of CTL lines specific for 5 different epitopes (SLYNTVATL, ILKEPVHGV, IEIKDTKEAL, SEGATPQDL, and KEKGGLEGL) to study the emergence of escape mutations. Escape varied between clones for the same epitope, and between different epitopes.

Gag and RT epitope escape, if it occurred at all, tended to be monoclonal and within the epitope, indicating strong fitness constraints, while the Nef epitope escape was rapid, polyconal, and sometimes the result of upstream frameshifts.

 There was one cloned cell line that recognized IEIKDTKEAL, 161JD27. After 2 weeks of passaging HIV-1 in the presence of 161JD27, no mutations were observed within the epitope in 10 sequences; one of the 10 had a single E -> K substitution 6 amino acids beyond the C-terminal end of the epitope.

HXB2 Location p17 (92-101)

**Author Location** Gag (92–101 B consensus)

Epitope IEIKDTKEAL

Epitope name IL10

Subtype B

Immunogen vaccine

Vector/Type: adeno-associated virus (AAV)

HIV component: gp120

Species (MHC) human (B60)

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay

Keywords dynamics, immune evasion

References Brainard et al. 2004

 HIV-1 gp120 is shown to suppress the ability of antigenspecific CTLs to migrate or remain at sites of high viral replication by concentration-dependent chemotaxis and fugetaxis. Directional T-cell movement is shown to depend on the interaction of the V2 and V3 loops with the CXCR4 receptor. X4 HIV-1 gp120 causes the migration of T-cells, including HIV-1 specific CTL, away from infected target cells, another potential mechanism for immune evasion.

**HXB2 Location** p17 (92–101)

Author Location p17 (92–101)

**Epitope** IEIKDTKEAL **Epitope name** Gag/p17-IL10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B60)

Assay type Chromium-release assay

Keywords binding affinity, epitope processing, TCR us-

age, characterizing CD8+ T cells

References Yang et al. 2003b

- Killing efficiency of CTLs and their ability to suppress viral replication is shown to depend more on epitope specificity than to antigenic avidity. Different clones recognizing the same epitope had similar killing efficiency despite their variation in avidity. Nef specific CTL clones tended to be most inhibitory, followed by Gag, then by RT specific clones, regardless of avidity.
- 1/14 CTL T-cell clones tested were specific for Gag/p17-IL10. Under conditions of excess peptide (100ug/ml), there was no difference in their lytic potential; all possessed similar effector capacity. Avidity was measured as the sensitizing dose of peptide required for 50% of maximal killing (SD50), which varied from 20 pg/ml to 100 ng/ml, over four orders of magnitude for all 14 epitopes. The SD50 value for the Gag/p17-IL10 clone was 8,000 pg/ml.

HXB2 Location p17 (92–101)

Author Location p17 (92–101)

Epitope IEIKDTKEAL

Immunogen HIV-1 infection

Species (MHC) human (B60, B61)

Keywords immunodominance

References Day et al. 2001

- No immunodominant responses were detected to five B61restricted epitopes tested.
- All five B60-restricted epitopes were reactive in another subject, and the B60-restricted responses together contributed over one-third of the total CTL response.

**HXB2 Location** p17 (93–101)

Author Location Gag (99-107 WEAU)

Epitope EVKDTKEAL

**Epitope name** Gag EVL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

**Donor MHC** A\*2902, B\*4403, B\*0801

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assav

Keywords dynamics, immunodominance, acute/early in-

fection, kinetics, characterizing CD8+ T cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance. WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.
- There was a weak response to this epitope during acute and early infection, and the epitope sequence did not vary during the first year of the infection.

**HXB2 Location** p17 (93–101)

**Author Location** p17

**Epitope** EVKDTKEAL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*0801)

**Donor MHC** A\*0101, A\*0301, B\*0801, B\*5101; A\*0101,

B\*0801

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , HLA binding

Keywords escape, acute/early infection, variant cross-

recognition or cross-neutralization

References Milicic et al. 2005

- CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient. In an early acute infection of the donor, the precise timing of transmission determines the viral variants transmitted.
- The second donor in the study shares A\*0101 and B\*0801 with his partner. The epitope EVKDTKEAL has an escape variant in the donor that does not react in an Elispot assay, DvkGtkeal, but the Dvkdtkeal form was the transmitted variant. The transmitted form and EVKDTKEAL bind with equal affinity to B\*0801.
- The recipient mounted a response to the Dvkdtkeal form of the epitope. The variant DvRdtkeal was detected by 32 weeks post infection.

HXB2 Location p17 (93–101) Author Location p17 (93–101) Epitope EIKDTKEAL

Epitope LINDINEAL

Immunogen peptide-HLA interaction

Species (MHC) human (B8)

References DiBrino et al. 1994b

Examined in the context of motifs important for HLA-B8 binding, predicted epitope based on Achour et al.

HXB2 Location p17 (93–101)
Author Location p17 (93–101)
Epitope EIKDTKEAL
Immunogen HIV-1 infection
Species (MHC) human (B8)
References Birk et al. 1998b

 A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

HXB2 Location p17 (93–101) Author Location p17 (93–101 LAI) Epitope EIKDTKEAL Subtype B

Immunogen

Species (MHC) human (B8, B60)

**References** Brander & Walker 1997

 Pers. comm. from A. Trocha and S. Kalams to C. Brander and B. Walker.

HXB2 Location p17 (93–101) Author Location p17 (SF2) Epitope DVKDTKEAL Immunogen HIV-1 infection Species (MHC) human

**Keywords** subtype comparisons, immunodominance

References Goulder et al. 2000a

 The CTL-dominant response was focused on this epitope in a HIV+ Caucasian from Boston, who was A1/\*0201 B8/63 Cw7/- – this epitope fell outside the most recognized peptides in the study.

- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

**HXB2 Location** p17 (119–127) **Author Location** Gag (119–127 BORI)

**Epitope** AADTGNSSQ **Epitope name** Gag AQ9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*2902, B\*1402, Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** dynamics, immunodominance, escape, acute/early infection, characterizing CD8+ T cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient BORI had high viral loads and rapid CD4 decline. BORI mounted 8 detected responses. BORI did not control viral replication well, and escape mutations occurred early and in most epitopes, 6/7 that were precisely identified.
- 20 variants in the AADTGNSSQ epitope were found in the patient BORI, the first appearing at day 35 with new variants continuing to arise through day 556. This is an extremely variable epitope, and changed not only by base substitution but by insertion and deletion. All variants tested conferred escape, at high concentrations of peptide.

HXB2 Location p17 (121–132)

Author Location p17 (121–132 HXB2R)

Epitope DTGHSNQVSQNY

Immunogen HIV-1 infection

Species (MHC) human (A33)

References Buseyne *et al.* 1993b

 Clustering of Gag p24 CTL epitopes recognized in 29 HIVinfected people.

HXB2 Location p17 (121–132) Author Location Gag (121–132 LAI) Epitope DTGHSNQVSQNY Subtype B

Immunogen HIV-1 infection Species (MHC) human (A33) References Buseyne *et al.* 1993a

- Vertical transmission of HIV ranges from 13% to 39%.
- Primary assays showed that cytotoxic activity against at least one HIV protein was detected in 70% of infected children.
- Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.
- Patient EM17 (CDC P2A+C+D2) had a CTL response to two epitopes in Gag.

**HXB2 Location** p17 (123–132)

**Author Location** Gag

Epitope GNSSQVSQNY

**Epitope name** GY10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 2, gKssqvsqny, was found not to correspond to the most polymorphic residues in the epitope.
   This is a novel partially mapped epitope.

**HXB2 Location** p17 (124–132)

**Author Location** p17 (124–132 LAI)

**Epitope NSSKVSQNY** 

Subtype B

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human (B\*3501)

**Keywords** optimal epitope

References Frahm et al. 2007

• Noted by Brander to be B\*3501 epitope.

**HXB2 Location** p17 (124–132)

**Author Location** p17

Epitope NSSQVSQNY Immunogen HIV-1 infection Species (MHC) human (B\*3501)

**Keywords** binding affinity **References** Dorrell *et al.* 2001

• The crystal structure of this epitope bound to HLA-B\*3501 shows that a serine can fit into the B pocket, which is shared between B35 and B53, with the hydroxyl group of the P2 serine occupying a position almost identical to the P2 proline that was previously considered the anchor motif.

Novel B53 epitopes (DTINEEAAEW and QATQEVKNM) were defined in this study that showed that A and T can also serve as P2 anchor residues for the B pocket of HLA-B35 and B53 – while S, T, and P could all fit into the B pocket and form a hydrogen bond, A would not form a bond, so the authors propose compensatory interactions account for the high affinity of QATQEVKNM for B53.

**HXB2 Location** p17 (124–132)

Author Location p17 (124–132 LAI)

**Epitope NSSKVSQNY** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords review

References McMichael & Walker 1994

• Review of HIV CTL epitopes.

**HXB2 Location** p17 (124–132)

**Author Location** 

Epitope NSSKVSQNY

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords dynamics, acute/early infection

References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers—high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

**HXB2 Location** p17 (124–132)

**Author Location** p17 (124–132)

Epitope NSSKVSQNY

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Birk et al. 1998b

 A study of p17 variation considering known p17 epitopes and individuals with known HLA types revealed that p17 evolution is influenced by immune pressure from CTLs.

**HXB2 Location** p17 (124–132)

Author Location p17 (124–132 LAI)

Epitope NSSKVSQNY

Subtype B

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human (B35)

References Rowland-Jones et al. 1995

• Established by titration.

HXB2 Location p17 (124–132) Author Location p17 (124–132 LAI) Epitope NSSKVSQNY

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (B35)

References Lalvani et al. 1997

- A peptide-based protocol was optimized for restimulation of CTLp using optimized peptide and IL-7 concentrations – importantly this protocol does not stimulate a primary response, only secondary – peptide-specific CTLp counts could be obtained via staining with peptide-Class I tetramers.
- This peptide was one of the B35 presented test peptides used in control experiments showing that the assay gave no activity using lymphocytes from 21 healthy B35 seronegative donors.

**HXB2 Location** p17 (124–132)

**Author Location** p17

**Epitope NSSKVSQNY** 

Immunogen

Species (MHC) human (B35)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied – these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive.
- HIV-2 version of this epitope is not conserved: PPSGKGGNY, but the CTLs are cross-reactive – this is one of five B35 CTL epitopes that are cross-reactive, see also Rowland-Jones *et al.* [1995].

**HXB2 Location** p17 (124–132)

**Author Location** p17

Epitope NSSKVSQNY Immunogen HIV-1 infection Species (MHC) human (B35) Keywords HAART, ART References Seth *et al.* 2001

 CTL responses were studied by tetramer staining in 41 patients with combination therapy – activated CD8+ T-cells decline as the viral load drops in response to therapy, but the overall level of antigen-specific cells capable of differentiating into effectors stays constant and new epitopes may be recognized.

**HXB2 Location** p17 (124–132)

Author Location p17 (124–132 SF2)

Epitope NSSKVSQNY
Immunogen HIV-1 infection
Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B35+ individuals that had a CTL response to this epitope broken down by group: 0/2 group 1, 0/2 group 2, and 1/1 group 3.

**HXB2 Location** p17 (124–132)

**Author Location** 

Epitope NSSKVSQNY

Epitope name Gag-NY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA B35, 1/21 (5%) recognized this epitope.

HXB2 Location p17 (124-132)

**Author Location** p17 (124–132)

**Epitope NSSKVSQNY** 

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2/9 patients recognized this epitope.

## II-B-2 Gag p17-p24 CTL/CD8 + epitopes

HXB2 Location p17-p24 (124-1)

Author Location Gag (124–133 BORI)

Epitope NSSQVSQNYP

Epitope name Gag NP10

Subtype B

Immunogen HIV-1 infection Species (MHC) human

Donor MHC A\*2902, B\*1402, Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

**Keywords** dynamics, immunodominance, escape, acute/early infection, characterizing CD8+ T cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient BORI had high viral loads and rapid CD4 decline.
   BORI mounted 8 detected responses. BORI did not control viral replication well, and escape mutations occurred early and in most epitopes, 6/7 that were precisely identified.
- 10 variants in the NSSQVSQNYP epitope were found in the patient BORI, the first appearing at day 35, NgSQVSQNYP, with new variants continuing to arise through day 556. This is an extremely variable epitope, and changed not only by base substitution but by insertion and deletion. All variants tested conferred some degree of escape by diminishing the CTL response.

**HXB2 Location** p17-p24 (126–11)

Author Location (C consensus)

Epitope GKVSQNYPIVQNLQGQMV

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B13) Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** p17-p24 (127-3)

**Author Location** p17-p24 (127–135 subtype D)

Epitope QVSQNYPIV

Subtype D

**Immunogen** 

Species (MHC) human (A\*6802)

References Dong 1998

- Epitope starts in p17 and ends in p24.
- · Predicted on binding motif, no truncations analyzed.

**HXB2 Location** p17-p24 (129–7)

Author Location Gag (129-139)

Epitope SQNYPIVQNIQ

**Epitope name** Gag 7.3 **Immunogen** vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT, Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining
Keywords subtype comparisons, variant crossrecognition or cross-neutralization, memory
cells

References Amara et al. 2005

- A clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better conserved for CD8 than CD4 T cells.
- All 5 CD8 and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation of a CD8 T cell epitope previously reported for humans as NYPIVQNL. HLA restriction: A\*2402.
- The response elicited to the B clade epitope SQNYPIVQNIQ does not cross-react with the CRF02 form SQNYPIVQNaQ. Other clades either most commonly carry an A or L in this position, SQNYPIVQN[a/l]Q.

**HXB2 Location** p17-p24 (131-6)

**Author Location** p17-p24 (132–140 SF2)

Epitope NYPIVQNL

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

References Ikeda-Moore et al. 1997

- The epitope starts in p17 and ends in p24.
- Defined using reverse immunogenetics 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) 53 of the 59 peptides bound A\*2402.
- This peptide induced CTL in 1/4 HIV-1 + people tested.
- NYPIVQNL bound to A\*2402 with medium strength, and the epitope can be processed in a vaccinia construct and presented no CTL clone was obtained.

## II-B-3 Gag p24 CTL/CD8 + epitopes

**HXB2 Location** p24 (3–11)

**Author Location** 

Epitope VQNLQGQMV

Epitope name VV9

Immunogen

Species (MHC) human (B13)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B13 epitope.

**HXB2 Location** p24 (8–17)

**Author Location** p24 (140–149)

Epitope GQMVHQAISP

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals was HLA A\*0201, A1, B57 and responded to four B57 epitopes and two others.

**HXB2 Location** p24 (8–20)

**Author Location** p24 (140–152 IIIB)

Epitope GQMVHQAISPRTL

Immunogen HIV-1 infection

Species (MHC) human (Cw3)

References Littaua et al. 1991

• Fine specificity of human Cw3 restricted Gag CTL epitope.

**HXB2 Location** p24 (8–20)

**Author Location** p24 (8–20)

Epitope GQMVHQAISPRTL

Immunogen HIV-1 infection

Species (MHC) human (Cw3)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these epitopes, but this increased over time.

**HXB2 Location** p24 (8–27)

**Author Location** p24 (140–159)

Epitope GQMVHQAISPRTLNAWVKVV

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Musey et al. 1997

• CTL specific for this epitope were found in the peripheral blood but not in the cervical mucosa of one donor.

**HXB2 Location** p24 (9–18)

**Author Location** Gag (173–182)

Epitope QMVHQAISPR

Immunogen HIV-1 infection

**Species (MHC)** human (A3 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

**HXB2 Location** p24 (9–23)

Author Location p24 (16-24)

Epitope QMVHQSLSPRTLNAW

Subtype A, D

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*3002, A\*6801, B\*5703, B\*5802; A\*3001,

A\*6601, B\*5801, B\*5802

Country Uganda.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, variant cross-

recognition or cross-neutralization

References Barugahare et al. 2005

- T-cell immune responses were examined in HIV-1 infected Ugandans. Similar levels of cross-clade responses for Gag, Env and Nef were observed. Higher frequencies of responses in conserved regions were found for Gag within a single clade, while areas with higher viral sequence variation had fewer detectable responses. Differential interclade sequence homologies between the Gag regions did not affect the respective level of cross-clade recognition.
- The sequence contains a known B7/B8 epitope, but the subjects recognizing it are B7- and B8-negative. The viral sequences isolated from the subjects were qmThqNlsprtlnaw and qmvhqAlsprtlnaw, and the peptide was recognized.

**HXB2 Location** p24 (10–18)

Author Location Gag (144–152 SF2)

Epitope MVHQAISPR

Subtype B

Immunogen HIV-1 infection, computer prediction

Species (MHC) human (A\*3303)

**Assay type** Chromium-release assay

**Keywords** binding affinity, computational epitope prediction

References Hossain et al. 2003

- HLA-A\*3303 is a common HLA allele in east and southeast Asia. Pol, Gag and Nef SF2 proteins were scanned for potential A\*3303 epitopes. 99 potential epitopes were synthesized, and 52/99 bound to A\*3303. Six of these served as peptide-targets for lysis by PBMC from infected individual, and clones derived from 4 of these 6 could lyse HIV-vaccinia infected target cells, indicating proper processing.
- This epitope is one of the 4 that are properly processed.

**HXB2 Location** p24 (10–18)

Author Location Gag (174–182)
Epitope MVHQAISPR
Immunogen HIV-1 infection
Species (MHC) human (A3 supertype)
Keywords supertype, rate of progression
References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 5/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location p24 (11–20) Author Location (C consensus) Epitope VHQAISPRTL

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*1510) Country South Africa.

**Assay type** CD8 T-cell Elispot - IFNγ **Keywords** characterizing CD8+ T cells **References** Kiepiela *et al.* 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location p24 (11–20) Author Location (C consensus) Epitope VH0AISPRTL

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*1510)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- VHQAISPRTL is an optimal epitope.

**HXB2 Location** p24 (11–24) **Author Location** p24 (SF2)

**Epitope** VQHAISPRTLNAWV **Immunogen** HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons, immunodominance **References** Goulder *et al.* 2000a

- The CTL-dominant response was focused on this epitope in an HIV+ Haitian living in Boston, who was A34/68 B57/71 Cw3/7 – this epitope fell outside the most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

HXB2 Location p24 (11-25)

**Author Location** p24 (11–25 HXB2)

Epitope VHQAISPRTLNAWVK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 29% of the study subjects, and it was the third most frequently recognized peptide.

HXB2 Location p24 (11–32)

Author Location p24 (143–164 BH10)

Epitope VHQAISPRTLNAWVKVVEEKAF

Immunogen HIV-1 infection Species (MHC) human (B57) References Johnson *et al.* 1991

• Gag CTL response studied in three individuals.

**HXB2 Location** p24 (12–20)

**Author Location** 

Epitope HQAISPRTL

Immunogen

Species (MHC) human (B\*1510)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes that this is an B\*1510 epitope.

HXB2 Location p24 (12–20)
Author Location Gag (146–154)
Epitope HQAISPRTL
Immunogen HIV-1 infection
Species (MHC) chimpanzee (Patr-B\*02)

**References** Balla-Jhagjhoorsingh *et al.* 1999b

- Certain HLA-alleles have been associated with long-term survival among them are HLA-B\*27 and HLA-B\*57.
- Of more than 150 chimpanzees that have been reported to be infected with HIV-1, only one has developed AIDS.
- CTL responses were studied in two HIV-1 infected chimpanzees that have strong CTL responses, and they were found to respond to highly conserved epitopes that are recognized in humans in the context of HLA-B\*27 and HLA-B\*57.
- The human HLA protein which presents this Patr-B\*02 epitope is HLA-B\*5701 but the amino acid sequences in the binding pockets of HLA-B\*5701 and Patr-B\*02 are distinctive.

**HXB2 Location** p24 (13–20)

**Author Location** p24 (145–152)

Epitope QAISPRTL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (Cw3)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

HXB2 Location p24 (13-23)

Author Location p24 (145–155 LAI)

Epitope QAISPRTLNAW

Subtype B

Immunogen

Species (MHC) human (A\*2501)

Keywords optimal epitope

**References** Frahm *et al.* 2007

• C. Brander notes that this is an A\*2501 epitope.

**HXB2 Location** p24 (13–23)

**Author Location** p24 (13–23 HXB2)

Epitope QAISPRTLNAW

Epitope name QW11

Subtype B

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Immunogen HIV-1 infection
Species (MHC) human (A*2501)
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**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** escape, immune evasion, optimal epitope, HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- Amino acid site in the third position potentially experienced positive selection. QAISPRTLNAW CTL escape mutant was found.

**HXB2 Location** p24 (13–23)

Author Location p24 (145–155 LAI)

Epitope QAISPRTLNAW

Subtype B

Immunogen

Species (MHC) human (A25)

References Kurane & West 1998

**HXB2** Location p24 (13–23)

**Author Location** p24 (145–155 SF2)

**Epitope** QAISPRTLNAV

Immunogen HIV-1 infection

Species (MHC) human (A25)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A25+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 0/2 group 2, and 1/3 group 3.

**HXB2 Location** p24 (13–23)

**Author Location** Gag (145–155 IIIB)

Epitope QAISPRTLNAW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A25)

Assay type Chromium-release assay

References Kurane et al. 2003

Three CD8+ CTL cell clones were derived from 2 HIV-1 positive asymptomatic patients, and their epitope specificities and HLA presenting proteins were defined.

HXB2 Location p24 (13–23)
Author Location p24 (145–155)
Epitope QAISPRTLNAW
Immunogen HIV-1 infection
Species (MHC) human

**Keywords** immunodominance **References** Betts *et al.* 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INFγ responses to other epitopes.
- 1/11 of the A2+ individuals was HLA A\*0201, A1, B57 and responded to QAISPRTLNAW noted previously to be A25.

**HXB2 Location** p24 (14–23)

**Author Location** Gag

**Epitope** AISPRTLNAW

Epitope name IW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Two escape mutations, at positions 1 (A146P) and 2 (I147L), were found in the most polymorphic residue in the epitope.
   Both were shared between clades B and C. Both were significantly more variable in persons expressing HLA-B57.

**HXB2 Location** p24 (14–23)

Author Location p24

Epitope AISPRTLNAW

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human (B57, B63)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

 HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and B58

 This epitope was recognized by 30% of B63-positive subjects and 29% of B57/58-positive subjects.

**HXB2 Location** p24 (15–23)

Author Location p24

Epitope LSPRTLNAW Immunogen HIV-1 infection Species (MHC) human (B\*57)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

**HXB2 Location** p24 (15–23)

**Author Location** p24 (147–155 IIIB)

Epitope ISPRTLNAW

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*5701 epitope.

**HXB2 Location** p24 (15–23)

**Author Location** 

**Epitope** ISPRTLNAW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords rate of progression, immunodominance

References Migueles & Connors 2001

HLA B\*5701 was found in a very high frequency in HIV-1 infected non-progressors, 11/13 (85%) versus 19/200 (9.5%) of progressors. Non-progressors tended to have an immune response that was highly focused on four p24 epitopes that were presented by B\*5701, ISPRTLNAW, KAFSPEVIPMF, TSTLQEQIGW, and QASQEVKNW.

HXB2 Location p24 (15-23)

**Author Location** 

Epitope ISPRTLNAW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

**Keywords** rate of progression, immunodominance

References Migueles & Connors 2001

- CTL activity was monitored in 27 individuals, including 10 LTNP with an over-expression of HLA B\*5701 – these individuals have viral loads below the threshold of infection without therapy, and their immune response tends to be focused on peptides that contain B\*5701 epitopes ISPRTLNAW, KAF-SPEVIPMF, TSTLQEQIGW, and QASQEVKNW.
- CTL responses are broader in B\*5701+ individuals with progressive viremia than those that control viremia.
- The HLA-A\*0201 SLYNTVATL epitope response was not as strong in individuals that carried both A2, B57.

HXB2 Location p24 (15-23)

**Author Location** 

Epitope ISPRTLNAW

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*5701)

**Assay type** Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

Keywords rate of progression, escape

References Migueles et al. 2003

- cDNA Gag sequences from a set of 17 HLA-B\*5701+ progressors and 10 LTNPs were obtained, and the variation in four p24 B\*5701 epitopes examined. Sequence variants were more common (p < 0.01) in the epitopes in the progressors (median 3, range 1-7) than LTNPs (median 2, range 0-4).
- In general use of the autologous protein in a target cell did not diminish the overall CD8+ T-cell responses.

**HXB2 Location** p24 (15–23)

**Author Location** Gag (147–155 LAI)

**Epitope** ISPRTLNAW

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5701, B\*5801)

**Keywords** rate of progression **References** Klein *et al.* 1998

- B57 has been associated with long-term non-progression in the Amsterdam cohort.
- The most pronounced CTL responses in HLA B\*5701 LTS were to RT and Gag.

**HXB2 Location** p24 (15–23)

Author Location p24 (15-23)

**Epitope** LSPRTLNAW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5801)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

 PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** p24 (15–23)

**Author Location** p24 (147–155)

Epitope ISPRTLNAW

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INFγ responses to other epitopes.
- 1/11 of the A2+ individuals was HLA A\*0201, A1, B57 and responded to four B57 epitopes and two others, but not SLYNT-VATI.

**HXB2 Location** p24 (15–23)

**Author Location** Gag (SF2)

Epitope ISPRTLNAW

Epitope name IW9

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords acute/early infection

References Goulder et al. 2001a

- This epitope elicited the second strongest CTL response in patient PI004 during acute infection, and maintained the response.
- Three CTL responses, to epitopes TSTLQEQIGW, ISPRTL-NAW, and KAFSPEVIPMF, were evident early after infection; CTL responses to SLYNTVATL, QASQEVKNW, EIYKRWII, and FLKEKGGL were detectable at 5 months post-infection and beyond.

HXB2 Location p24 (15-23)

**Author Location** p24 (147–155)

Epitope ISPRTLNAW

Epitope name ISP

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- None of the 8 study subjects recognized this epitope but none were HLA B57+.

**HXB2 Location** p24 (15–23)

Author Location p24 (15-23)

Epitope ISPRTLNAW

Immunogen HIV-1 infection Species (MHC) human (B57) References Ferrari *et al.* 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location p24 (15–23)
Author Location p24 (147–155 SF2)
Epitope ISPRTLNAW
Immunogen HIV-1 infection
Species (MHC) human (B57)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B57+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 0/0 group 2, and 2/2 group 3.

**HXB2 Location** p24 (15–23)

**Author Location** 

**Epitope** ISPRTLNAW **Epitope name** Gag-IW9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

References Sabbaj et al. 2003

- Among HIV+ individuals who carried HLA B57, 2/5 (40%) recognized this epitope.
- Among HIV+ individuals who carried HLA B58, 0/4 (0%) recognized this epitope.

**HXB2 Location** p24 (15–23)

**Author Location** 

**Epitope ISPRTLNAW** 

**Epitope name** ISP

Immunogen HIV-1 infection Species (MHC) human (B57)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius *et al.* [2000, 2001a] in an IFN? Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there was no correlation between CTL responses with viral rebound rates, plateau viral loads, or clearance rates.

Author Location p24 (15–23)
Author Location Gag (147–155)
Epitope ISPRTLNAW
Immunogen HIV-1 infection
Species (MHC) human (B57)

**Donor MHC** A3, A28, B53, B57; A31, B7, B57

Assay type Chromium-release assay

**Keywords** TCR usage, genital and mucosal immunity

References Musey et al. 2003

- CTL clones from blood, semen, cervix and rectum samples from 12 individuals were compared. CTL clones derived from blood and mucosal samples had similar high lysis efficiency, primarily perforin dependent, and TCR? VDJ region sequencing revealed cases of expansion of the same clone in different compartments.
- CD8+ T cell clones directed at this epitope were derived from blood and semen of one male subject, and blood and cervix of one female subject.
- From the male patient, six clones that recognized this epitope had three different patterns of TCR? usage: 2 from the blood and 1 from the semen used V $\beta$ 6S2DJ2S2; 1 from the blood and 1 from the semen used V $\beta$ 6S2DJ1.1; and 1 from the semen used V $\beta$ 7S1DJ2.3.
- From the female patient, five clones that recognized this epitope had different TCR? usage. Blood derived clones were B?6S7DJ2.7, B?6.4DJ2.3, and B?6S3DJ2.1. Cervix derived clones were B?6S3DJ1.4 and B?6S5DJ2.5.

HXB2 Location p24 (15–23) Author Location Gag (147–155) Epitope ISPRTLNAW

Epitope name ISW9 Subtype B, C

Immunogen HIV-1 infection Species (MHC) human (B57)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

Keywords epitope processing, escape

References Draenert et al. 2004b

• The A146P mutation flanking the ISW9 epitope (Pisprtlnaw) is positively selected in HLA-B57+ persons and it prevents trimming of the optimal epitope by the endoplasmic reticulum aminopeptidase I. The A146P processing escape mutation does not influence replicative capacity of the virus in vitro and is accumulated over time in the human population.

**HXB2 Location** p24 (15–23)

**Author Location** p24 (15–23)

Epitope ISPRTLNAW

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 3/7 patients recognized this epitope.

**HXB2 Location** p24 (15–23)

Author Location (147–155 B consensus)

Epitope ISPRTLNAW

Epitope name IW9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords characterizing CD8+ T cells

References Allen et al. 2004

- This study characterizes an escape mutation in a C-terminal flanking residue of the HLA-A3 gag p17 KK9 epitope that inhibits processing, and is embedded in the overlapping HLA-A3 RK9 epitope.
- The immune response was tracked in subject AC-38. The acute immunodominant response was to the B57 TW10 epitope; this response declined following viral escape (tsNlqeqigw) by day 64. The p17 KK9 and RK9 became immunodominant, but then declined as the escape mutation arose. IW9 is 1 of 3 other strong responses that persisted, along with 1 sub-dominant response.

**HXB2 Location** p24 (15–23)

Author Location p24

Epitope ISPRTLNAW

**Epitope name** ISW9

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords review, epitope processing, escape

References Goulder & Watkins 2004

This paper is a review of the role of CTL in HIV infection, and it
uses the ISW9 epitope as an example of an epitope that escapes
due to a mutation before the N-terminal end of the epitope.
The insertion of a proline prevents the aminopeptidase ERAAP
from cleaving the glutamine from the precursor, qPisrptlnaw,
preventing processing of ISRPTLNAW.

**HXB2 Location** p24 (15–23)

**Author Location** Gag

Epitope ISPRTLNAW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, epitope processing, es-

cape, characterizing CD8+ T cells, reversion,

viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Characteristic changes in B57 epitopes in B57+ people were mapped: ISPRTLNAW often has the substitution Lsprtlnaw, as well as the proximal A->P substitution Pisprtlnaw.

**HXB2 Location** p24 (15–23)

**Author Location** p24 (147–155)

Epitope ISPRTLNAW

Epitope name IW9

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Ethiopia.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords immunodominance, escape, variant cross-

recognition or cross-neutralization

References Currier et al. 2005

- Epitope sequence variation and CD8 T-cell responses were analyzed in C subtype infected HLA-B57-positive individuals from Ethiopia. KF11 was the immunodominant response.
- ISPRTLNAW had a variant ISPRTLNAW in 7/10 B57+ subjects, and 4/9 B57- subjects; 2 other variants were observed, but there was no apparent sequence selection in this epitope.

**HXB2 Location** p24 (15–23)

**Author Location** Gag (147–155)

**Epitope** LSPRTLNAW

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Canada.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay

Keywords mimics

References Mason et al. 2005

- CTL responses against the human IP-30 signal peptide sequence LLDVPTAAV were shown to be elicited by stimulation of PBMCs from HIV-1 infected individuals with HIV protease peptide 76-84, LVGPTPVNI. In vitro stimulation with HIV PR 76-84 or the IP-30 signal peptide was shown to activate a comparable population of cross-reactive effector cells. None of the peptides activated CTL in non-HIV-infected individuals. IP-30 signal peptide was shown to have lower avidity T-cell interactions than the HIV peptide.
- As a control, responses to A2-restricted HIV epitopes ALVE-ICTEM, EELRQHLLRW, and LSPRTLNAW were shown not to give IP-30 responses.

**HXB2 Location** p24 (15–23)

Author Location Gag (147–155)

Epitope ISPRTLNAW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

**Donor MHC** A\*3001, A\*66, B\*4201, B\*5802, Cw\*0602, Cw\*1701: A\*68, A\*66, B\*57, B\*5802.

Cw\*1/01; A\*68, A\*66, B\*5/, B\*5802 Cw\*0602, Cw\*0701

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** epitope processing, responses in children, mother-to-infant transmission, escape, acute/early infection

References Pillay et al. 2005

- HIV-specific CTLs in infants were shown to be able to select for viral escape variants early in life, despite a lack of escape with the same CTL specificity in the mother. Infant CTL responses may be compromised by transmission of escape variants that arose in the mother and also those that arose in the father, if the father was the source of the mother's infection.
- ISPRTLNAW is the C consensus form of the epitope and was the autologous form in the mother, and was transmitted to her infant. By 33 weeks a new dominant form of the epitope had emerged in the infant, mSPRTLNAW, and two additional variants had arisen, one with a substitution proximal to the epitope, plSPRTLNAW, and lSPRTLNAW.

**HXB2 Location** p24 (15–23)

**Author Location** p24 (147–155 IIIB)

Epitope ISPRTLNAW
Immunogen HIV-1 infection
Species (MHC) human (B57, B\*5801)
Keywords rate of progression
References Goulder et al. 1996b

- Five slow progressors made a response to this epitope, and in two it was the dominant response.
- Peptide defined on the basis of B\*5801 binding motif, yet not cross-restricted except at high concentrations.

**HXB2 Location** p24 (15–23)

**Author Location** Gag

**Epitope ISPRTLNAW** 

**Epitope name** ISW9

Immunogen HIV-1 infection Species (MHC) human (B57, B\*5801)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** epitope processing, responses in children, mother-to-infant transmission, escape

References Feeney et al. 2005

- Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.
- ISPRTLNAW was recognized more often in children than in adults, and was the most frequently recognized B57 epitope in children. Escape variants of this epitope arose in 2 children: an A->P change proximal to the epitope, pISPRTLNAW, and and

I1L change, ISPRTLNAW. In both cases the mother carried AISPRTLNAW.

**HXB2 Location** p24 (15-23)

**Author Location** p24 (subtype A)

Epitope LSPRTLNAW

Subtype A

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B57, B58)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix systemic CD8+ T cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

**HXB2 Location** p24 (15–23)

Author Location p24 (147–155)

**Epitope** LSPRTLNAW

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B57, B58)

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

- Variants (L/I)SPRTLNAW are specific for the A/B clades.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused
  on different epitopes with HLA presenting molecules that have
  previously been associated with reduced risk of infection, and
  there was a shift in the response in the HEPS women upon late
  seroconversion to epitopes recognized by the HIV-1 infected
  women
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B57/B58 women, 4/6 HEPS and 14/17 HIV-1 infected women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in 2 of the 4/6 HEPS cases and in 7 of the 14/17 responsive HIV-1 infected women.

**HXB2 Location** p24 (15–23)

**Author Location** p24

**Epitope** ISPRTLNAW

Subtype B, D

Immunogen HIV-1 infection

Species (MHC) human (B58)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

## References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence had an IIL change, ISPRTLNAW.

**HXB2 Location** p24 (15–23)

**Author Location** 

**Epitope** ISPRTLNAW

Immunogen

Species (MHC) human (B63)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B63 epitope.

**HXB2 Location** p24 (15–23)

**Author Location** 

**Epitope** KAFSPEVIPMF

Immunogen

Species (MHC) human (B63)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B63 epitope.

**HXB2 Location** p24 (15–23)

**Author Location** Gag

Epitope ISPRTLNAW

Subtype B

Immunogen HIV-1 infection Species (MHC) human (Cw\*0602)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- I->L (Lsprtlnaw) is associated with HLA C\*0602.

HXB2 Location p24 (15-23)

**Author Location** 

Epitope LSPRTLNAW

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- ISPRTLNAW was consistently recognized by 1/22 HEPS sex worker controls (ML1250), and LSPRTLNAW was recognized by 2 additional HEPS sex worker controls (ML1693 and ML1589).

**HXB2 Location** p24 (16–24)

**Author Location** p24 (148–156)

**Epitope** SPRTLNAWV

Immunogen

Species (MHC) human (B\*0702)

Keywords optimal epitope

References Frahm et al. 2007

- C. Brander notes this is a B\*0702 epitope.
- Optimal peptide mapped by titration.

**HXB2 Location** p24 (16–24)

Author Location p24 (148–156)

Epitope SPRTLNAWV

Immunogen

Species (MHC) human (B7)

References Brander & Walker 1997

 Optimal peptide mapped by titration, pers. comm. from D. Lewinsohn to C. Brander and B. Walker.

HXB2 Location p24 (16-24)

**Author Location** p24 (148–156)

Epitope SPRTLNAWV

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Brodie et al. 2000

- Study tracks and quantifies in vivo migration of neo-marked CD8 HIV-specific CTL.
- Adoptively transferred gene-marked HIV-specific CTL homed to specific lymph node sites, colocalizing within the parafollicular regions of the lymph node adjacent to cells expressing HIV tat-fusion transcripts, indicative of viral replication.
- The CTL clones expressed CCR5 and localized among HIV-1 infected cells expressing MIP-1alpha and MIP-1beta, CCchemokines produced at sites of viral replication, suggesting a possible homing mechanism.
- This study provides a methodology for tracking and studying antigen specific CTL *in vivo*.

**HXB2 Location** p24 (16–24)

Author Location p24 (148–156)

**Epitope** SPRTLNAWV

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B7)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Subject ML 1203 started with CTL responses to A\*6802 DTVLEDINL and to B7 FPVTPQVPLR prior to seroconversion, and upon seroconversion acquired additional responses to A\*6802 ETAYFILKL which became dominant, B7 TPGPGVIRYPL, B7 IPRRIRQGL, and B7 SPRTLNAWV.

HXB2 Location p24 (16-24)

Author Location p24 (16-24)

**Epitope** SPRTLNAWV

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP).
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

HXB2 Location p24 (16-24)

**Author Location** 

**Epitope** SPRTLNAWV

Epitope name Gag-SW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Sabbaj et al. 2003

- Among HIV+ individuals who carried HLA B07, 1/9 (11%) Only 20% of CD8+ T-cells produce IFN-gamma and granzyme recognized this epitope.
- Among HIV+ individuals who carried HLA B81, 1/6 (17%) recognized this epitope.

**HXB2 Location** p24 (16–24)

Author Location p24 (16-24)

**Epitope** SPRTLNAWV

Epitope name B7-SV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

Keywords dynamics, supervised treatment interruptions

(STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 1/11 HLA-B7 positive individuals had detectable B7-restricted responses to this epitope during acute infection - 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 1/4 individuals had detectable responses to this epitope after STI.

**HXB2 Location** p24 (16–24)

Author Location p24 (16-24)

Epitope SPRTLNAWV

Epitope name B7-SV9 Gag

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFNγ

Keywords supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

· An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response, but in this case the epitope did not

**HXB2 Location** p24 (16–24)

Author Location p24 (148–156)

**Epitope** SPRTLNAWV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, CD8 T-cell Elispot granzyme B

Keywords characterizing CD8+ T cells

References Kleen et al. 2004

B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.

- None of seven patients responded to this peptide with GzB producing cells or IFN-gamma producing cells.
- The authors describe the epitope as SPRTLNNQWV the double N's may be a typo or an unusual form of the epitope; it is atypical and may be why there was no response.

**HXB2 Location** p24 (16–24) **Author Location** p24 (16–24)

Epitope SPRTLNAWV Immunogen HIV-1 infection Species (MHC) human (B7)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 7 patients recognized this epitope.

**HXB2 Location** p24 (16–24)

Author Location p24 (subtype B)

**Epitope** SPRTLNAWV

Subtype B

**Immunogen** HIV-1 exposed seronegative

Species (MHC) human (B7, B\*8101)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix systemic CD8+ T cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

**HXB2 Location** p24 (16–24)

**Author Location** Gag (subtype B)

**Epitope** SPRTLNAWV

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B7, B\*8101)

Keywords subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among A, B, and D clade viruses.

**HXB2 Location** p24 (16–24)

Author Location p24

**Epitope** SPRTLNAWV

Immunogen HIV-1 infection

Species (MHC) chimpanzee

References Santra et al. 1999

- 3/4 animals displayed HIV-1 Gag-specific CTL activity.
- Effector cells from two chimpanzees were able to recognize epitopes also recognized by human HIV-1 Gag-specific CTL (SPRTLNAWV, HLA-B7, and DLNTMLNTV, HLA-B14)
- No chimpanzee CTL were detected to the following human HIV-1 specific Gag epitopes, although they were embedded within 20mer peptides that contained a reactive epitope: ISPRTLNAW, HLA-B57; KRWIILGLNK, HLA-B27; and DRFYKTLRA, HLA-B14.

**HXB2 Location** p24 (16–25)

**Author Location** Rev

Epitope QIRSLSGWIL

Epitope name QL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A2, B7, B44, Cw5, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 7, QIRSLSeWIL was found not to correspond to the most polymorphic residue in the epitope. This is a novel unmapped epitope.

**HXB2 Location** p24 (18–26)

**Author Location** Gag (150–)

Epitope RTLNAWVKV

**Epitope name** Gag150

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: p24 Gag Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human, transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

 HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient. • This peptide was an intermediate A2 binder, and induced CTL responses in mice. Responses were detected in 1/17 HIV+ HLA-A2 subjects.

HXB2 Location p24 (19-26)

**Author Location** Gag

**Epitope** TLNAWVKW

Epitope name T9V

Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV compo-

nent: gp140, gp140∆V3

Species (MHC) transgenic mouse (A\*0201)

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted
CTL responses against multiple conserved HIV-1 epitopes in
HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw
virus expressing gp140 with deleted V1, V2, and V3 loops
successfully induces neutralizing antibodies against HIV-1. A
live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination
against HIV and measles.

**HXB2 Location** p24 (19–27)

Author Location p24 (151–159)

Epitope TLNAWVKVV

Immunogen HIV-1 infection

**Species (MHC)** human (A\*02)

Keywords HAART, ART, immunodominance

References Huang et al. 2000

- The single cell ELISPOT assay was optimized and highly specific, and found to work well even after the primary cells had been frozen and thawed.
- Increases in gamma interferon producing cells were observed in response to anti-retroviral therapy using single cell IFNgamma-production ELISPOT.
- In 3/3 HLA-A\*02, -B\*27 subjects the immunodominant epitope was against HLA B\*27 Gag p24 epitope KRWIILGL, not A2 Gag epitopes.

**HXB2 Location** p24 (19–27)

Author Location p24 (151–159)

Epitope TLNAWVKVV

Immunogen HIV-1 infection

**Species (MHC)** human (A\*02)

Keywords HAART, ART

References Rinaldo et al. 2000

Administration of triple-drug antiretroviral therapy (IDV, 3TC and ZDV) sometimes showed a transient increase and other times failed to increase CTL responses in patients with advanced HIV disease, but there is a stable population of tetramer stained HIV-specific CD8+ CD45RO+ cells that can persist after therapy and long periods of virus being below the level of detection.

HXB2 Location p24 (19-27)

Author Location p24 (151–159)

Epitope TLNAWVKVV

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Parker et al. 1992; Parker et al. 1994

 Study of sequence motifs preferred for peptide binding to class I HLA-A2.

**HXB2 Location** p24 (19–27)

**Author Location** p24 (19–27)

**Epitope** TLNAWVKVV

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (19–27)

**Author Location** p24 (150–159)

Epitope TLNAWVKVI

References Kaul et al. 2001a

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A2)

**Keywords** subtype comparisons, HIV exposed persis-

tently seronegative (HEPS)

- Variants TLNAWVKV(I/V) are A/B clade specific.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** p24 (19–27)

**Author Location** p24 (A02, A30, B4402, B15)

Epitope TLNAWVKVV

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining

Keywords HIV exposed persistently seronegative

(HEPS), characterizing CD8+ T cells

References Missale et al. 2004

- HIV-specific T-cell response was tested in patients exposed to blood from a patient with highly replicating HIV; these patients were nosocomially infected with HBV, but uninfected with HIV. HIV-specific T-cell responses were directed to structural and non-structural HIV proteins in 2 patients, suggesting that the virus replicated in these patients sufficiently to prime a cell-mediated immune response that protected them from HIV infection.
- This patient responded to 4/8 HIV epitopes tested in an IFNgamma EliSpot assay or tetramer assay. Responses were detected to this peptide 8 and 28 weeks after exposure with EliSpot, but not by tetramer binding.

**HXB2 Location** p24 (19–27)

**Author Location** p24 (19–27 HXB2)

Epitope TLNAWVKVI

Epitope name 24D

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: multiple epitope immunogen HIV component: p17/p24 Gag,

Pol Adjuvant: IL-12

**Species (MHC)** transgenic mouse (A2)

Assay type Cytokine production, CD8 T-cell Elispot -IFNγ, Chromium-release assay

Keywords vaccine-specific epitope characteristics, vaccine antigen design

References Bolesta et al. 2005

- Immunization of transgenic mice with a codon-optimized hGagp17p24-Polp51 DNA plasmid, consisting of clusters of highly conserved CTL epitopes presented by multiple MHC class I alleles, induced 2- to 5-fold higher CD8+ T-cell responses than the corresponding full-length proteins. The modified proteins had the ribosomal frameshift deleted, as well as the potentially immunosuppressive p15, and protease and integrase. This correlated with higher protection against challenge with Gag and Pol expressing recombinant vaccinia virus. Mice immunized with the hGagp17p24-Polp51 also showed an elevated level of type 1 cytokine production as well as an increased titer of p24- and RT-specific IgG2 antibody responses.
- This was 1 of 4 A2 gag/pol epitopes tested. Transgenic mice immunized with the deleted construct induced more potent EliSpot reactions to this epitope than those immunized with full length Gag/Pol.

HXB2 Location p24 (19-27)

**Author Location** p24 (subtype B)

**Epitope** TLNAWVKVV

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2, A\*0202)

Keywords subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi - these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among A, B and D clade viruses.

**HXB2 Location** p24 (21–30)

**Author Location** Gag (153–162 WEAU)

**Epitope** NAWVKIEEK Epitope name Gag NK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A\*2902, B\*4403, B\*0801

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords dynamics, immunodominance, acute/early in-

fection, kinetics, characterizing CD8+ T cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTLmediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance. WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.
- There was a weak response to this epitope during acute and early infection, and the epitope sequence did not vary during the first year of the infection.

HXB2 Location p24 (21-31)

Author Location Gag (129-139)

Epitope NAWVKVVEEKA

Epitope name Gag 8.4 Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT,

Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining

**Keywords** subtype comparisons, memory cells

References Amara et al. 2005

- A clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better conserved for CD8 than CD4 T cells.
- All 5 CD8 and 4/8 of the CD4 epitopes that were recognized in vaccinated macagues have also been reported for human infections, indicating cross-species conservation. The similar reported human epitope in this case is NAWVKVVEEKAF-SPEVIPMF, HLA restriction: A2, B21, B57.
- The B clade immune response to NAWVKVVEEKA gives a diminished response to the CRF02 variant NAWVKViEEKA, but does cross-react. The M group clades are about evenly split between the 2 variants.

**HXB2 Location** p24 (21–40)

Author Location Gag (153–172)

Epitope NAWVKVVEEKAFSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Brodie et al. 1999

• The ability of CTL effector cells was studied by expanding autologous HIV-1 Gag-specific CTL in vitro, and adoptively transferring them.

The transferred CTLs migrated to the lymph nodes and transiently reduced circulating productively infected CD4+ T cells, showing that CTL move to appropriate target sites and mediate anti-viral effects.

**HXB2 Location** p24 (21–40) **Author Location** p24 (153–172)

Epitope NAWVKVVEEKAFSPEVIPMF

Immunogen HIV-1 infection
Species (MHC) human (B57)
References Brodie et al. 2000

- Study tracks and quantifies in vivo migration of neo-marked CD8+ HIV-specific CTL.
- Adoptively transferred gene-marked HIV-specific CTL homed to specific lymph node sites, colocalizing within the parafollicular regions of the lymph node adjacent to cells expressing HIV tat-fusion transcripts, indicative of viral replication.
- The CTL clones expressed CCR5 and localized among HIV-1 infected cells expressing MIP-1? and MIP-1?, CC-chemokines produced at sites of viral replication, suggesting a possible homing mechanism.
- This study provides a methodology for tracking and studying antigen specific CTL *in vivo*.

**HXB2 Location** p24 (21–40)

Author Location p24 (153-172 SF2)

Epitope NAWVKVVEEKAFSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A2, -B21.

**HXB2 Location** p24 (21–40)

Author Location p24 (153–172 SF2)

Epitope NAWVKVVEEKAFSPEVIPMF

Immunogen vaccine

Vector/Type: virus-like particle (VLP) HIV

component: CD4BS, Gag, gp120, V3

Species (MHC) macaque

References Wagner et al. 1998b

- A VLP is a non-infectious virus-like particle self-assembled from HIV Pr55 gag macaques were immunized with VLPs bound to either gp120 or V3+CD4 linear domains Gag and Env specific CTL were stimulated in each case, and Ab response to gag and gp120 was elicited, but the gp120 neutralizing response occurred only with whole gp120, not V3+CD4 despite the CTL and Ab response, immunized macaques were infected by intervenous challenge with SHIV chimeric challenge stock Wagner *et al.* [1998b]
- CTL specific for this epitope could be found both before and after SHIV challenge.

**HXB2 Location** p24 (21–42)

**Author Location** p24 (153–174 BH10)

Epitope NAWVKVVEEKAFSPEVIPMFSA

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Johnson et al. 1991

• Gag CTL response studied in three individuals.

**HXB2 Location** p24 (24–32)

**Author Location** (C consensus)

Epitope VKVIEEKAF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (24–32)

**Author Location** (C consensus)

Epitope VKVIEEKAF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- VKVIEEKAF is an optimal epitope.

**HXB2 Location** p24 (24–32)

**Author Location** 

Epitope VKVIEEKAF

Immunogen

Species (MHC) human (B\*1503)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B\*1503 epitope.

**HXB2 Location** p24 (27–36)

Author Location p24 (27–36)

**Epitope** IEEKAFSPEV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4006)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** p24 (27–37)

**Author Location** (C consensus)

Epitope IEEKAFSPEVI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4501)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- IEEKAFSPEVI is an optimal epitope.

**HXB2 Location** p24 (28–36)

Author Location p24

Epitope EEKAFSPEV

Subtype A

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*4415)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Bird et al. 2002

- 5/233, (4 HIV-1 positive, 1 HEPS) (2.1%) Kenyan female sex workers carried the novel HLA allele B\*4415.
- Residues forming the B pocket of HLA B\*4415 were identical to HLA B\*4001, B\*4402 and B\*4403. These alleles preferred E, an acidic residue, at the P2 position.
- The amino acid residues forming the F pocket of allele B\*4415 were not correlated with other known HLA molecules, but analogy suggests a binding preference for small, neutral amino acids.
- Based on the binding motif x[DE]xxxxxx[VILA], 19 potential B\*4415 epitopes were identified, and 1/19 was reactive in an Elispot, EEKAFSPEV.

HXB2 Location p24 (28-36)

Author Location p24 (28-36)

Epitope EEKAFSPEV

Immunogen

Species (MHC) human (B\*4415)

**Keywords** optimal epitope

References Frahm et al. 2007

**HXB2 Location** p24 (28–36)

Author Location (C consensus)

**Epitope** EEKAFSPEV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4501)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (28–47)

**Author Location** p24 (160–179)

Epitope EEKAFSPEVIPMFSALSEGA

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Musey et al. 1997

 Cervical and peripheral blood derived CTL clones from an HIV-infected woman recognized this epitope.

HXB2 Location p24 (29–36)

Author Location p24 (28–36)

Epitope EKAFSPEV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0602)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

 $\label{lem:keywords} \textbf{Keywords} \ \ \text{subtype comparisons, computational epitope}$ 

prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** p24 (29–39)

Author Location Gag (161–171)

Epitope EKAFSPEVIPM

Epitope name Gag 8.5

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT,

Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining
Keywords subtype comparisons, variant crossrecognition or cross-neutralization, memory
cells

References Amara et al. 2005

- A clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better conserved for CD8 than CD4 T cells.
- All 5 CD8 and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation. The similar reported human epitope in this case is EEKAFSPEVIPMF-SALSEGA, HLA restriction: B27
- This epitope is conserved in all HIV-1 clades except CRF01, and is identical in B and CRF02.

HXB2 Location p24 (29-48)

**Author Location** Gag (161–180 C consensus)

**Epitope** EKAFSPEVPMFTALSEGAT

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** p24 (30–37)

Author Location p24 (162–170 LAI)

 ${\bf Epitope} \ {\sf KAFSPEVI}$ 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5703)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*5703 epitope.

**HXB2 Location** p24 (30–37)

Author Location p24 (30–37)

**Epitope** KAFSPEVI

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Goulder et al. 2000c

- Two strong clonal CTL responses were generated in donor 026-BMC (HLA A3/-, B42/B57, Cw7/17) against different optimal versions of this epitope, one 8 amino acids long, one 11.
- Improved stabilization of the B57-peptide complex was demonstrated by the 11 mer which fits the B57 binding motif, relative to the 8 mer, which does not.
- B57 tolerates marked difference in optimal peptide length and B57 is associated with non-progressive infection.

**HXB2 Location** p24 (30–37)

**Author Location** 

**Epitope** KAFSPEVI

Epitope name Gag-KI8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Sabbaj et al. 2003

 Among HIV+ individuals tested who carried HLA B57, 0/5 (0%) recognized this epitope.

**HXB2 Location** p24 (30–40)

Author Location p24

Epitope KAFSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human (B\*57)

Keywords HAART, ART

References Spiegel et al. 1999

- Study examines the effect of highly active antiretroviral therapy (HAART) on HIV-1 plasma viral load, CTLp and CTLe frequencies in 8 infected children.
- CTLp (precursors) were measured by stimulating in culture and assaying using 51Cr release, against vaccina expressed IIIB Env, Gag, Pol, Nef, and CTLe were measured by ELISPOT.
- CTL against B\*57-KAFSPEVIPMF was a de novo response observed in one of the children when viral load increased as a result of stopping therapy.
- HIV-1 specific CTL responses initially increased in children with complete viral suppression, but then decreased, suggesting viral replication is needed to maintain CTL responses.

**HXB2 Location** p24 (30–40)

**Author Location** Gag

**Epitope** KAFSPEVIPMF

Epitope name KF11

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human (B\*57)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, HLA binding

**Keywords** subtype comparisons, escape, reversion, viral fitness, optimal epitope

**References** Leslie *et al.* 2005

 An escape mutation, A2G (KgFSPEVIPMF), is suggested to be a result of selection pressure from the HLA-B\*57 allele, and can be transmitted and stable in the absence of HLA-B\*57. Evidence indicated that the mechanism of escape was an increased off-rate.

**HXB2 Location** p24 (30–40)

Author Location p24 (162–172 LAI)

Epitope KAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords rate of progression

References Goulder et al. 1996b

- This peptide was recognized by CTL from five slow progressors.
- Peptide defined on the basis of B\*5801 binding motif, yet not cross-restricted except at high concentrations.
- This epitope is highly conserved.

**HXB2 Location** p24 (30–40)

Author Location p24 (162–172 LAI)

Epitope KAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords optimal epitope

**References** Frahm *et al.* 2007

• C. Brander notes this is a B\*5701 epitope.

**HXB2 Location** p24 (30–40)

**Author Location** 

**Epitope** KAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5701)

Keywords rate of progression, immunodominance

**References** Migueles & Connors 2001

- HLA B\*5701 was found in a very high frequency in HIV-1 infected non-progressors, 11/13 (85%) versus 19/200 (9.5%) of progressors. Non-progressors tended to have an immune response that was highly focused on four p24 epitopes that were presented by B\*5701, ISPRTLNAW, KAFSPEVIPMF, TSTLQEQIGW, and QASQEVKNW.
- Attempts to make all for HLA B\*5701-epitope tetramers were made, but only the HLA B\*5701-KAFSPEVIPMF tetramer folded properly. The percentage of CD8+ T cells staining with this HLA B\*57 gag tetramer and the fraction of CD69+IFN-+ cells responding to autologous B cells pulsed with KAFSPE-VIPMF was highly correlated (r = 0.84; P = 0.005). The percent of CD8+ T cells that stain with the A\*2 gag SLYNTVATL tetramer was low (0-0.31%) in a A2+ B57+ LTNP, emphasizing the focus of the immune response on the B\*5701 epitopes.

**HXB2 Location** p24 (30–40)

**Author Location** 

Epitope KAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords rate of progression, immunodominance

References Migueles & Connors 2001

 CTL activity was monitored in 27 individuals, including 10 LTNP with an over-expression of HLA B\*5701 – these individuals have viral loads below the threshold of infection without therapy, and their immune response tends to be focused on peptides that contain B\*5701 epitopes ISPRTLNAW, KAFSPEVIPMF, TSTLQEQIGW, and QASQEVKNW.

- CTL responses are broader in B\*5701+ individuals with progressive viremia than those that control viremia.
- The HLA-A\*0201 SLYNTVATL epitope response was not as strong in individuals that carried both A2 and B57.

**HXB2 Location** p24 (30–40)

Author Location Gag (162–172)

**Epitope** KAFSPEVIPMF

Epitope name KAF11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Assay type Intracellular cytokine staining, Flow cytomet-

ric T-cell cytokine assay

Keywords rate of progression, escape

References Migueles et al. 2003

- cDNA Gag sequences from a set of 17 HLA-B\*5701+ progressors and 10 LTNPs were obtained, and the variation in four p24 B\*5701 epitopes examined. Sequence variants were more common (p < 0.01) in the epitopes in the progressors (median 3, range 1-7) than LTNPs (median 2, range 0-4).</li>
- In general, use of the autologous protein in a target cell did not diminish the overall CD8+ T-cell responses.
- This epitope tends to be quantitatively immunodominant in B57+ people, including in some of the individuals in this study.
   It was extremely well conserved in the sequences obtained here, despite strong immune pressure, suggesting fitness constraints.

**HXB2 Location** p24 (30–40)

**Author Location** p24 (30–40)

Epitope KAFSPEVIPMF

**Epitope name** KAFS

Subtype A, B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5701, B\*5703)

**Keywords** subtype comparisons, rate of progression

References Gillespie et al. 2002

- CTL responses of eight HIV+ slow progressors from Nairobi Kenya or Oxford, UK who were B\*5701 or B\*5703 were studied, as B\*57 is associated with slow progression.
- This epitope is located between the structurally conserved alpha-helix 1 and alpha-helix 2 (H1-H2) region of the p24 capsid protein, and tends to elicit strong reactions in B\*57 individuals.
- Broad heterogeneous cross-clade reactivity to 6 clade variants of the KAFS peptide sequence were observed in one B\*5701 and 5 B\*5703 HLA-restricted patients, measured by IFNγ production Elispot assays as well as tetramer binding. The clade variants were: KAFSPEVIPMF (clades A and B), kGfNpevipmf (clades A/AC); kaLspevipmf (clade A); kafspevipVf (clade A); kafNpeIipmf (group O); kafspeIipmf (A/C); kafsQevipmf (A/C); and kaLspevipmf KNFSPEVIPMF A/G). Not all variants were well recognized in all patients, for example kafsQevipmf was not able to induce IFN gamma production in 3/6 tested, and had a diminished capacity to sensitize target cells for lysis.

**HXB2 Location** p24 (30–40)

Author Location p24 (162-172 LAI)

Epitope KAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5703)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is a B\*5703 epitope.

HXB2 Location p24 (30-40)

**Author Location** 

**Epitope** KAFSPEVIPMF

Epitope name Gag-KF11

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*5703)

**Donor MHC** A\*3402 A\*7401 B\*0801 B\*5703 Cw\*0302

Cw\*0701

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a
- Subject 00RCH59 was African American, on HAART, viral load 170, CD4 count 477.
- recognized this epitope.

**HXB2 Location** p24 (30–40)

**Author Location** p24 (162–172)

**Epitope** KAFSPEVIPMF

Epitope name KF11

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5703)

Country Ethiopia.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, immunodominance, es-

cape, variant cross-recognition or crossneutralization

References Currier et al. 2005

- HLA-B57 is associated with slow progression. Epitope sequence variation and CD8 T-cell responses were analyzed in HLA-B\*5703-positive individuals from Ethiopia. KF11 epitope and its variants were found to be immunodominant in these subjects. Two HLA-B\*5702 subjects did not respond to the KF11 epitope or its variants.
- 5 variants of the epitope were observed: KAFSPEVIPMF, Kn-FSPEVIPMF, rAFSPEVIPMF, KgFnPEVIPMF, and KAFSPE-VIPMI. Depending on the subject, different versions of these variants were more or less susceptible to their CD8+ T cells, i.e., one person's escape form was another person's susceptible form.

HXB2 Location p24 (30-40) Author Location (C consensus)

Epitope KAFSPEVIPMF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5703)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the A2 and S4 residues of KAFSPE-VIPMF are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** p24 (30–40)

Author Location p24 (30–40)

**Epitope** KAFSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Goulder et al. 2000c

- · Two strong clonal CTL responses were generated in donor 026-BMC (HLA A3/-, B42/B57, Cw7/17) against different optimal versions of this epitope, one 8 amino acids long, one 11.
- Improved stabilization of the B57-peptide complex was demonstrated by the 11mer which fits the B57 binding motif, relative to the 8 mer, which does not.
- Among HIV+ individuals who carried HLA-B57, 6/6 (100%) B57 tolerates marked difference in optimal peptide length and B57 is associated with non-progressive infection.

**HXB2 Location** p24 (30–40)

**Author Location** p24 (162–172)

**Epitope** KAFSPEVIPMF

**Immunogen** HIV-1 infection

Species (MHC) human (B57)

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals was HLA A\*0201, A1, B57 and responded to four B57 epitopes and two others.

HXB2 Location p24 (30-40)

**Author Location** p24 (SF2)

**Epitope** KAFSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

• The CTL-dominant response was focused on this epitope in a HIV+ Caucasian living in Boston – this epitope is not among the most recognized peptides in the study.

- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

**HXB2 Location** p24 (30–40) Author Location Gag (SF2) **Epitope** KAFSPEVIPMF

Epitope name KF11

Immunogen HIV-1 infection Species (MHC) human (B57) References Goulder et al. 2001a

• Three CTL responses in patient PI004, to epitopes TSTLQE-QIGW, ISPRTLNAW, and KAFSPEVIPMF, were evident early after infection; CTL responses to SLYNTVATL, QASQEVKNW, EIYKRWII, and FLKEKGGL were detectable

HXB2 Location p24 (30-40) **Author Location** p24 (162–172) **Epitope** KAFSPEVIPMF **Epitope name** KAF

at 5 months post-infection and beyond.

Immunogen HIV-1 infection Species (MHC) human (B57)

Kevwords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- None of the 8 study subjects recognized this epitope but none were HLA B57+.

HXB2 Location p24 (30-40)

Author Location p24

**Epitope** KAFSPEVIPMF Immunogen HIV-1 infection Species (MHC) human (B57)

References Kostense et al. 2001

- HLA tetramers to six epitopes were used to study HLA-A2, B8 and B57 CTL in 54 patients - HIV-specific tetramer positive cells were inversely correlated with viral load in patients with high CD4, but in patients with CD4 T-cells below 400 high tetramer frequencies were found despite high viral load.
- · Most patients have high levels of HIV-specific T-cell expansions, but many of these cells aren't functional.
- In 15 of the patients, the proportion of IFN gamma producing tetramer cells correlated with AIDS-free survival.

**HXB2 Location** p24 (30–40)

**Author Location** p24 (162–172 SF2)

Epitope KAFSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B57+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 0/0 group 2, and 2/2 group 3.

**HXB2 Location** p24 (30–40)

Author Location p24 (163–174)

Epitope KAFSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Appay et al. 2000

- · Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virusspecific CD8+ T cells produced IFN- $\gamma$  and MIP-1 $\beta$  with a distinct subset that failed to produce TNF- $\alpha$ .

**HXB2 Location** p24 (30–40)

**Author Location** 

**Epitope** KAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA B57, 1/5 (20%) recognized this epitope.

HXB2 Location p24 (30-40)

**Author Location** p24

**Epitope** KAFSPEVIPMF

Epitope name KAF

Immunogen HIV-1 infection Species (MHC) human (B57)

Keywords HAART, ART, supervised treatment interrup-

tions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius et al. [2000, 2001a] in an IFN? Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there
  was no correlation between CTL responses with viral rebound
  rates, plateau viral loads, or clearance rates.

HXB2 Location p24 (30–40) Author Location p24 (30–40) Epitope KAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

**Donor MHC** A\*0201, A3, B44, B57, Cw5, Cw6

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** acute/early infection, early-expressed proteins **References** Cao *et al.* 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- Alleles A3, B35, B57, and B62 were more frequently recognized than alleles A1, A2, A30, and A44 e.g., during primary infection. 2/10 patients, 1372 and 1397, recognized A2-restricted epitopes. The common A2-restricted epitopes Gag SL9 and Pol IV9 were not recognized in peptide tetramer-binding assays.

**HXB2 Location** p24 (30–40)

Author Location p24

Epitope KAFSPEVIPMF Immunogen HIV-1 infection Species (MHC) human (B57)

Assay type Intracellular cytokine staining

**Keywords** immunodominance, genital and mucosal immunity

References Kaul et al. 2003

- Predefined immunodominant peptide responses were used to compare CD8+ T-cell responses in the blood and cervix of 16 HIV+ Kenyan sex workers. Cervical responses were detected in 8/10 women from whom adequate samples could be obtained. The frequency of the CD8+ T-cell response in the genital tract was comparable to the blood, with a trend toward being slightly higher.
- The immunodominant response was to this epitope in the PBMC of 10/16 patients (Kaul et al. 2001, AIDS, 107:1303).

HXB2 Location p24 (30–40) Author Location p24 (163–174) Epitope KAFSPEVIPMF Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

**Donor MHC** A\*0201, A3, B57, Cw\*06, Cw\*07; A\*01, A\*0201, B\*08, B\*57, Cw6, Cw7

Country United States.

Assay type Cytokine production, Tetramer binding, Intracellular cytokine staining, Degranulation, CD107a and b cell surface mobilization

**Keywords** TCR usage **References** Betts *et al.* 2004

• Both cytokine production and degranulation in HIV-1 specific and CMV specific CD8+ T -cells occurs at high peptide concentrations together with TCR downregulation. Only degranulation is observed at lower peptide concentrations with no observed TCR downregulation. Thus the nature of CTL response depends not on the specific T cell clonotype or antigen, but on the concentration of Ag presented on APCs.

**HXB2 Location** p24 (30–40)

Author Location p24

Epitope KAFSPEVIPMF

Epitope name TW10

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** epitope processing, escape **References** Draenert *et al.* 2004b

 This study characterizes the N-terminal flanking position of the epitope ISPRTLNAW, and mutations in this position are thought to impact processing. The B57 epitope KAFSPE-VIPMF was used as a positive control in this study.

**HXB2 Location** p24 (30–40)

Author Location Gag (155–172 B con)

Epitope KAFSPEVIPMF

Epitope name KF11

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords variant cross-recognition or crossneutralization

References Draenert et al. 2004c

• CD8+ T-cell responses that persisted in individuals with advanced disease were studied; most of the optimal epitopes defined were recognized with intermediate to high avidity. On average 13 (range, 2-39) epitopic regions were targeted in an average of 6 proteins (range, 1-8). HAART resulted in decrease in antigen and reduction in gamma IFN EliSpot responses, suggesting active responses to autologous virus. The lack of mutations within most viral epitopes suggest that persistent CTL through late disease do not exert strong immune selection

pressure, yet the Elispot assays show robust responses, suggesting to the authors that gamma IFN-based screening methods may not reveal functional CD8+ T-cell impairment in patients with AIDS.

2 subjects recognized this epitope, 1 with high functional avidity, 1 with intermediate. Autologous sequence revealed no substitutions in this epitope compared to the B consensus.

HXB2 Location p24 (30-40)

**Author Location** Gag

**Epitope** KAFSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** HIV exposed persistently seronegative (HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- 1/2 HLA B57+ infection-resistant men, compared to 0/1 preseroconversion men who went on to become infected, reacted to this epitope.

**HXB2 Location** p24 (30–40)

**Author Location** p24 (30–40)

Epitope KAFSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI)

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 7/7 patients recognized this epitope.

**HXB2 Location** p24 (30–40)

Author Location (162–172 B consensus)

Epitope KAFSPEVIPMF

Epitope name KF11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

**Keywords** immunodominance, characterizing CD8+ T cells

References Allen et al. 2004

- This study characterizes an escape mutation in a C-terminal flanking residue of the HLA-A3 gag p17 KK9 epitope that inhibits processing, and is embedded in the overlapping HLA-A3 RK9 epitope.
- The immune response was tracked in subject AC-38. The acute immunodominant response was to the B57 TW10 epitope; this response declined following viral escape (tsNlqeqigw) by day 64. The p17 KK9 and RK9 became immunodominant, but then declined as the escape mutation arose. FK11 is 1 of 3 other strong responses that persisted, along with 1 sub-dominant response.

**HXB2 Location** p24 (30–40)

Author Location (C consensus)

Epitope KAFSPEVIPMF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location p24 (30-40)

Author Location p24

Epitope KAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country United Kingdom.

Assay type Tetramer binding, T-cell Elispot, Intracellular

cytokine staining

**Keywords** rate of progression, acute/early infection, characterizing CD8+ T cells, immune dysfunction

References Papagno et al. 2004

Acute HIV-1 infection induces massive activation of HIV-specific and non-HIV-specific CD8+ T-cells resulting in differentiation of these cells. High differentiation of CD8+ T-cells is correlated with disease progression. Differentiation is a natural process but it can be driven by elevated immune activation, such as in HIV infection.

**HXB2 Location** p24 (30–40)

**Author Location** Gag

Epitope KAFSPEVIPMF

Epitope name KF11 Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was quite conserved in people carrying B57, but two substitutions were found in 11 B57+ individuals tested: kNfspevipmf and kafspelipmf.

**HXB2 Location** p24 (30–40)

Author Location Gag (162-172)

**Epitope** KAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** p24 (30–40)

Author Location Gag (162-172 BRU)

**Epitope** KAFSPEVIPMF **Subtype** B, CRF02\_AG **Immunogen** HIV-1 infection

Species (MHC) human (B57)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons **References** Inwoley *et al.* 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 1/9 CRF02\_AG-infected Ivorians, and 1/9 B-infected French subjects.

**HXB2 Location** p24 (30–40)

**Author Location** Gag

Epitope KAFSPEVIPMF

Epitope name KAF

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Netherlands.

Assay type Tetramer binding, Flow cytometric T-cell cy-

tokine assay

Keywords binding affinity, rate of progression, escape,

characterizing CD8+ T cells

References Jansen et al. 2005

- HLA-B57 has been associated with long term non-progression in HIV+ people. The number and responsiveness of CD8 T-cells directed to different Gag peptides presented by HLA-A2, -B8 and B57 were compared. T cells specific for the HLA-B57 epitope KAFSPEVIPMF responded to a higher extent and more readily to antigenic stimulation than those specific for the A2 epitope SLYNTVATL and the B8 epitope EIYKRWII.
- Tetramer decay experiments indicate that the HLA-B57 peptide has a higher half-life than the A2 and B8 peptides. The authors point out that CD8+ T cells with high binding affinity may require less help.
- Variant forms of the HLA-B57 epitope KAFSPEVIPMF were found in the 3/5 HLA B57+ individuals sequenced, but the variants were always a minor form.

**HXB2 Location** p24 (30–40)

**Author Location** 

Epitope KAFSPEVIPMF Immunogen HIV-1 infection Species (MHC) human (B57, B\*5801)

Aggregation CD9 T call Eligant 1ET

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** responses in children, mother-to-infant transmission, escape, characterizing CD8+ T cells

References Feeney et al. 2005

- Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.
- KAFSPEVIPMF was frequently recognized in children and in

**HXB2 Location** p24 (30–40)

**Author Location** p24 (153–164)

Epitope KAFSPEVIPMF

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (B57, B58)

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B57/B58 women, 4/6 HEPS and 12/17 HIV-1 infected women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in 2 of the 4/6 HEPS cases and in 7 of the 12/17 HIV-1 infected women.

HXB2 Location p24 (30–40) Author Location p24 (30–40) Epitope KAFSPEVIPMF Immunogen HIV-1 infection Species (MHC) human (B57, B58)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

**HXB2 Location** p24 (30–40)

**Author Location** p24

**Epitope** KAFSPEVIPMF

Subtype B, C

Immunogen HIV-1 infection Species (MHC) human (B57, B63)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This epitope was recognized by 60% of B63-positive subjects and 33% of B57/58-positive subjects.

HXB2 Location p24 (30–40) Author Location p24 (30–40) Epitope KAFSPEVIPMF Immunogen HIV-1 infection Species (MHC) human (B58)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (30–40)

**Author Location** p24

Epitope KAFSPEVIPMF

Subtype B, G

Immunogen HIV-1 infection

Species (MHC) human (B58)

Donor MHC A2, A36, B45, B58, Cw3, Cw6

Country Nigeria.

**Assay type** CD8 T-cell Elispot - IFNγ **Keywords** subtype comparisons

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype G Gag. The autologous epitope sequence in this person matched the known epitope.

HXB2 Location p24 (30-40)

**Author Location** 

Epitope KAVRLIKFLY

Immunogen

Species (MHC) human (B63)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes that this is an B63 epitope.

**HXB2 Location** p24 (30–40)

Author Location p24

Epitope KAFSPEVIPMF

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized by 1/22 HEPS sex worker controls, ML1250.

**HXB2 Location** p24 (31–44)

Author Location p24 (31–44 HXB2)

**Epitope** AFSPEVIPMFSALS

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided, it appears to be HXB2.
- Responses to this peptide were detected in 22% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p24 (31–50)

**Author Location** p24 (163–182)

Epitope AFSPEVIPMFSALSEGATPQ

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

• HIV-specific CTL lines developed by *ex vivo* stimulation with peptide.

**HXB2 Location** p24 (31–50)

**Author Location** p24 (163–182 SF2)

Epitope AFSPEVIPMFSALSEGATPQ

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A2, B21.

**HXB2 Location** p24 (31–50)

Author Location p24 (163–182 SF2)

Epitope AFSPEVIPMFSALSEGATPQ

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

 CTL expanded ex vivo were later infused into HIV-1 infected patients.

**HXB2 Location** p24 (31–50)

Author Location p24 (SF2)

Epitope AFSPEVIPMFSALSEGATPQ

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.
- The response to the peptide was CD4 dependent, but the HLA presenting molecule and optimal epitope were not determined.

**HXB2 Location** p24 (32–40)

**Author Location** Gag (164–172)

Epitope FSPEVIPMF

Immunogen HIV-1 infection

Species (MHC) human (B57)

**Donor MHC** A3, A28, B53, B57

Assay type Chromium-release assay

Keywords TCR usage, genital and mucosal immunity

References Musey et al. 2003

- CTL clones from blood, semen, cervix and rectum samples from 12 individuals were compared. CTL clones derived from blood and mucosal samples had similar high lysis efficiency, primarily perforin dependent, and TCR? VDJ region sequencing revealed cases of expansion of the same clone in different compartments.
- CD8+ T cell clones directed at this epitope were derived from blood and semen.
- The TCR? VDJ rearrangement of a CTL clone from the blood was V $\beta$ 21S3DJ1.2, and a clone from the semen used V $\beta$ 7S1DJ2.3.

HXB2 Location p24 (32-40)

**Author Location** 

Epitope FSPEVIPMF

Immunogen

Species (MHC) human (B57)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B57 epitope.

**HXB2 Location** p24 (32–40)

Author Location p24

Epitope FSPEVIPMF

Epitope name FF9

Subtype B, C

Immunogen HIV-1 infection

**Species (MHC)** human (B63, B57, B58)

**Donor MHC** A\*74, A\*8001, B\*18, B\*57, Cw\*02, Cw\*07

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords cross-presentation by different HLA, optimal

epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This is a putative HLA-B63/57/58 epitope containing the B58 supertype binding motif. Significantly more often recognized by B63+ and B57/58+ subjects than by negative subjects. Optimal epitope defined.

**HXB2 Location** p24 (33–40) **Author Location** p24 (33–40)

**Epitope** SPEVIPMF

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*35)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

HXB2 Location p24 (34–49) Author Location p24

Epitope PEVIPMFSALSEGATP

Immunogen in vitro stimulation or selection

Species (MHC)

Assay type Other

Keywords assay standardization/improvement, charac-

terizing CD8+ T cells

References Stone et al. 2005

 A new microarray technique was developed to screen small samples of T cells for specific peptide-MHC binding and functional responses. Each array element acts as an artificial antigen-presenting cell, consisting of immobilized recombinant MHC-peptide complex, costimulatory molecules, and cytokine-capture antibodies. The elements specifically elicit T-cell responses such as adhesion, secretion of cytokines, and modulation of surface markers.

HXB2 Location p24 (35–43) Author Location p24 (167–175 LAI) Epitope EVIPMFSAL

> Subtype B Immunogen

Species (MHC) human (A\*2601)

Keywords subtype comparisons

References Goulder et al. 1996a

- Identified as optimal epitope within Gag sequence AFSPE-VIPMFSALSEGATPQ.
- Relatively conserved epitope within B clade and in other clades.
- Suspected binding motif for HLA-A26 includes T or V anchor at position 2, negative charge at position 1.
- C. Brander notes that this is an A\*2601 epitope in the 1999 database.

**HXB2 Location** p24 (35–43)

Author Location p24 (167-175 LAI)

**Epitope** EVIPMFSAL

Subtype B

Immunogen

Species (MHC) human (A\*2601)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an A\*2601.

HXB2 Location p24 (35-43)

**Author Location** Gag (169–177)

**Epitope** EVIPMFSAL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2601)

Country Japan.

Assay type Chromium-release assay, Flow cytometric Tcell cytokine assay, Other, HLA binding

**Keywords** subtype comparisons, immunodominance, optimal epitope

References Satoh et al. 2005

- Reverse immunogenetics was used to identify HIV-1 epitopes presented by HLA-A\*2601. 110 peptides were predicted to bind to HLA-A\*2601. 24 of these were demonstrated to bind through a HLA-A\*2601 stabilization assay. Four of these, including this one, were shown to be epitopes endogenously presented by this allele, that can induce peptide-specific CD8 T-cells. HLA-A\*2601 is common in Asia.
- Immunodominant epitope recognized in 5/7 HIV-infected individuals with HLA-A\*2601. This epitope is highly conserved in clade B and E (CRF01).

**HXB2 Location** p24 (35–43)

Author Location (C consensus)

**Epitope** EVIPMFTAL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2601)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- EVIPMFTAL is an optimal epitope.

**HXB2 Location** p24 (35–43)

Author Location Gag (169–177 SF2)

**Epitope** EVIPMFSAL

Subtype A, B, C, CRF01\_AE, D

Immunogen HIV-1 infection

**Species (MHC)** human (A\*2601, A\*2603)

Country Japan.

Assay type Chromium-release assay, Flow cytometric Tcell cytokine assay, HLA binding

Keywords binding affinity, subtype comparisons, rate of progression, immunodominance, escape, cross-presentation by different HLA, variant cross-recognition or cross-neutralization

References Kawashima et al. 2005

- A\*26 is associated with slow progression to disease and is common in Asian populations (about 20%). 31/110 HIV peptides that carried the A\*2603 motif ([VTILP] at P2, [ML] at the C-terminus) bound to HLA-A\*2603. Only 2 of these were epitopes and could induce specific CD8 T-cell responses in PBMC from HLA-A\*2603 positive subjects.
- This epitope induced specific CD8+ T cells in chronically infected individuals with either A\*2603 or A\*2601. It is an immunodominant epitope.
- 3 common B clade variants were synthesized. EVIPMFaAL and EVIPMFtAL bound to A\*2603 with equal affinity as the consensus form, EVIPMFSAL, but could not be recognized by an EVIPMFSAL-specific T-cell clone so may mediate TCR escape. The other common variant, kVIPMFSAL, did not bind to A\*2603.
- EVIPMFSAL is the most common form in clades A, B, D, and E (CRF01), but EVIPMFtAL is the most common form in clade C.

HXB2 Location p24 (35-43)

Author Location p24 (167–175)

Epitope EVIPMFSAL

Immunogen HIV-1 infection

Species (MHC) human (A26)

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals that didn't respond to SLYNTVATL reacted with seven other epitopes including this epitope.

HXB2 Location p24 (35-43)

**Author Location** Gag

Epitope EVIPMFSAL

Epitope name EL9

Immunogen HIV-1 infection

Species (MHC) human (A26)

Donor MHC A26, B27

Assay type CD8 T-cell Elispot - IFNγ

Keywords responses in children, rate of progression, im-

munodominance, escape

References Feeney et al. 2004

• Viral load in a perinatally infected child remained low until emergence of an escape variant (kTwiilglnk) in the immunodominant CTL epitope KRWIILGLNK when the child was 7.4 years old. The emergence of this escape mutation was followed by an increase in viremia and an increase in the number of targeted CTL epitopes, measured again when the child was 9.2 years old. The EL9 response was not observed until after the escape mutation occurred in the immunodominant epitope, and was detected in the 9.2 year sample for the first time.

**HXB2 Location** p24 (35–49)

**Author Location** p24 (35–48 HXB2)

**Epitope** EVIPMFSALSEGATP

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- Responses to this peptide were detected in 16% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p24 (36–43)

**Author Location** p24 (168–175 LAI)

**Epitope VIPMFSAL** 

Subtype B

Immunogen

Species (MHC) human (Cw\*0102)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a C\*0102(Cw1) epitope.

**HXB2 Location** p24 (36–43)

Author Location p24 (36–43 HXB2)

**Epitope VIPMFSAL** 

Epitope name VL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0102)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101, Cw\*0102, Cw\*1203

Assav type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, optimal epitope, HIV-1

References Liu et al. 2006

• T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.

**HXB2 Location** p24 (36–43) Author Location p24 (168–175 LAI) **Epitope VIPMFSAL** Subtype B

Immunogen

Species (MHC) human (Cw\*0102, Cw1)

References Goulder et al. 1997b

**HXB2 Location** p24 (36–43) **Author Location** p24 (168–175) **Epitope** VIPMFSAL Immunogen HIV-1 infection Species (MHC) human (Cw1, Cw2)

Keywords immunodominance References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immuno-
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals that didn't respond to SLYNTVATL reacted with seven other epitopes including this epitope.

**HXB2 Location** p24 (37–52)

Author Location Gag (169–184 LAI)

Epitope IPMFSALSEGATPQDL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B12)

References Buseyne et al. 1993a

- Vertical transmission of HIV ranges from 13% to 39%
- Primary assays showed that cytotoxic activity against at least one HIV protein was detected in 70% of infected children.
- Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.
- Patient EM17 (CDC P2A+C+D2) had a CTL response to two epitopes in Gag.

**HXB2 Location** p24 (37–52)

**Author Location** p24 (169–184 LAI)

**Epitope** IPMFSALSEGATPQDL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B44)

References Buseyne et al. 1993b

• Clustering of Gag p24 CTL epitopes recognized in 29 HIVinfected people.

**HXB2 Location** p24 (37–52)

Author Location p24 (37-52)

Epitope IPMFSALSEGATPDQL

Immunogen HIV-1 infection

Species (MHC) human (B44)

References Ferrari et al. 2000

One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (39–58)

**Author Location** Gag (171–190)

**Epitope MFTALSEGTPQDLNTMLNT** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 8 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location p24 (41–60)

Author Location p24 (179–188 subtype A)

Epitope SALSEGATPQDLNMMLNIVG

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (B\*8101)

Keywords subtype comparisons

References Dorrell et al. 1999

- CTL responses in three individuals with non-clade B infections were studied, 2 with subtype A infections, 1 with subtype C their infections all originated in East Africa.
- This CTL epitope is presented by B\*8101 in one of the patients with an A subtype infection - B\*8101 is a newly discovered HLA allele found in Africans, and the epitope has yet to be mapped precisely.
- This epitope is distinct in subtype A relative to subtypes B, C, and D which share the dominant sequence: SALSEGATPQDL-NTMLNTVG.

**HXB2 Location** p24 (41–60)

Author Location p24 (173–192 SF2)

Epitope SALSEGATPQDLNTMLNTVG

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- Three of these 12 had CTL response to this peptide.

• The responding subjects were HLA-A3, A32, B7, B14; and HLA-A2, A3, B14, B44.

• The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct

**HXB2 Location** p24 (41–60)

Author Location p24 (173-192 SF2)

Epitope SALSEGATPQDLNTMLNTVG

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

 CTL expanded ex vivo were later infused into HIV-1 infected patients.

**HXB2 Location** p24 (41–60)

Author Location p24 (SF2)

Epitope SALSEGATPQDLNTMLNTVG

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.
- The response to the peptide was CD8 dependent, but the HLA presenting molecule and optimal epitope were not determined.

**HXB2 Location** p24 (41–62)

**Author Location** p24 (173–194 BH10)

Epitope SALSEGATPQDLNTMLNTVGGH

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Johnson et al. 1991

• Gag CTL response studied in three individuals.

**HXB2 Location** p24 (43–52)

**Author Location** Gag (175–184 WEAU)

Epitope LSEGATPQDL

Epitope name Gag LL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*4403)

**Donor MHC** A\*2902, B\*4403, B\*0801

Country United States.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

Keywords dynamics, immunodominance, escape, kinet-

ics, characterizing CD8+ T cells, reversion,

viral fitness

References Jones et al. 2004

• Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.

- The patient WEAU had high viral loads and rapid CD4 decline.
   WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance.
   WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.
- There was a weak response to this epitope during acute and early infection, and the epitope sequence did not vary during the first year of the infection.

**HXB2 Location** p24 (43–52)

**Author Location** p24 (subtype A)

**Epitope** LSEGATPQDL

Subtype A

**Immunogen** HIV-1 infection

Species (MHC) human (B42, B44)

Keywords subtype comparisons

References Cao et al. 2000

- HIV-1 subtypes A and D dominate the Ugandan epidemic, and a vaccine trial using B clade antigen is underway – this study addresses relative levels of cross-reactive CTL responses in Ugandans to A, D, and B clade recombinant vaccinia viruses expressing Gag, Env, Pol, RT or Nef from HIV-1 clades A, B, and D.
- Proteins corresponding to the subtype of the infecting strains tended to trigger higher levels of CTL response measured by percent specific lysis, but there was extensive inter-subtype cross-reactivity with B clade proteins and the co-circulating subtype.
- This optimal epitope sequence, recognized by CTL derived from a Ugandan with an A subtype infection (patient SP 511), is cross-reactive with subtypes A, B and D peptides.

**HXB2 Location** p24 (44–52)

**Author Location** p24 (176–184)

Epitope SEGATPQDL

Immunogen

Species (MHC) human (B\*4001)

Keywords optimal epitope

**References** Frahm *et al.* 2007

• C. Brander notes this is a B\*4001, B60 epitope.

**HXB2 Location** p24 (44–52)

**Author Location** Gag (178–186 BRU)

Epitope SEGATPQDL

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (B44)

Country Cote D'Ivoire.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 1/9 CRF02\_AG-infected Ivorians, and 1/9 B-infected French subjects.

**HXB2 Location** p24 (44–52) Author Location p24 (SF2) **Epitope** SEGATPQDL Immunogen HIV-1 infection Species (MHC) human (B60) References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes.
- B60 is present in 10-20% of the Caucasoid and very common in Asian populations.

HXB2 Location p24 (44-52)

**Author Location** p24

Epitope SEGATPQDL

Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (B60)

**Donor MHC** A2, A24, B38, B60, Cw2, Cw12

Assay type CD8 T-cell Elispot - IFNγ

Keywords HAART, ART, supervised treatment interrup-

tions (STI), acute/early infection, early treat-

References Montefiori et al. 2003

• HIV-1 + patient AC10 underwent early HAART treatment, which was discontinued 1.5 years later. At this timepoint potent NAb responses against autologous virus were detected. Treatment interruption initially induced weak CD8+ responses directed against 5 epitopes. By days 873d and 923d the CTL response had broadened to target 22 epitopes; of these six were fully characterized. Eventually the virus escaped the NAb response, but escape was not accompanied by a rise in viral load, and the authors suggest the virus was contained by the CTL response.

**HXB2 Location** p24 (44–52)

Author Location p24 (44-52 NL43)

**Epitope** SEGATPQDL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B60)

Assay type Chromium-release assay, CTL suppression of replication

Keywords escape

References Yang et al. 2003a

- Virus was cultured in the presence of CTL lines specific for 5 different epitopes (SLYNTVATL, ILKEPVHGV, IEIKDTKEAL, SEGATPQDL, and KEKGGLEGL) to study the emergence of escape mutations. Escape varied between clones for the same epitope, and between different epitopes. Gag and RT epitope escape, if it occurred at all, tended to be monoclonal and within the epitope, indicating strong fitness constraints, while the Nef epitope escape was rapid, polyconal, and sometimes the result of upstream frameshifts.
- One CTL clone, 161Jx12, recognized this epitope, and apparently no resistance mutations were selected by this clone, although the data was not shown in the paper.

HXB2 Location p24 (44-52) **Author Location** p24 (176–184) Epitope SEGATPQDL

Epitope name Gag/p24-SL9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B60)

**Assay type** Chromium-release assay

Keywords binding affinity, TCR usage, characterizing CD8+ T cells

References Yang et al. 2003b

- Killing efficiency of CTLs and their ability to suppress viral replication is shown to depend more on epitope specificity than to antigenic avidity. Different clones recognizing the same epitope had similar killing efficiency despite their variation in avidity. Nef specific CTL clones tended to be most inhibitory, followed by Gag, then by RT specific clones, regardless of avidity.
- 1/14 CTL T-cell clones tested were specific for Gag/p24-SL9. Under conditions of excess peptide (100ug/ml), there was no difference in their lytic potential; all possessed similar effector capacity. Avidity was measured as the sensitizing dose of peptide required for 50% of maximal killing (SD50), which varied from 20 pg/ml to 100 ng/ml, over four orders of magnitude for all 14 epitopes. The SD50 value for Gag/p24-SL9 was 30 pg/ml, it was among the peptides with the highest avidity.

**HXB2 Location** p24 (44–52)

**Author Location** p24 (HXB2)

Epitope SEGATPQDL

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (B60)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords epitope processing, vaccine antigen design, characterizing CD8+ T cells

References SenGupta et al. 2004

• Multiple HLA calss I-restricted and class II-restricted T-cell epitopes were shown to be processed and presented from an exogenously added HIV-1 gag-p24 peptide complexed to a heat shock protein. T-cell recognition of the complex was shown to be inhibited by brefeldin A indicating an endoplasmic reticulum-dependent pathway.

**HXB2 Location** p24 (44–52)

**Author Location** p24 (44–52)

Epitope SEGATPQDL

Immunogen HIV-1 infection

Species (MHC) human (B60, B61)

Keywords immunodominance

References Day et al. 2001

- No immunodominant responses were detected to five B61restricted epitopes tested.
- All five B60-restricted epitopes were reactive in another subject, the strongest CTL response directed against the B60-epitope p24 SEGATPQDL, and the B60-restricted responses together contributed over one-third of the total CTL response.

HXB2 Location p24 (46–59)

Author Location p24 (SF2)

**Epitope** GATPQDLNTMLNTV Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons, immunodominance References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in a HIV+ African American living in Boston with HLA A\*3002/68 B14/\*5802 Cw6/8 – this epitope fell within the most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects from South Africa.

HXB2 Location p24 (47-55) Author Location p24 (47-55) Epitope ATPQDLNTM Immunogen HIV-1 infection Species (MHC) human (B7) References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2** Location p24 (47–56) **Author Location** p24 (subtype A) **Epitope** ATPQDLNMML Subtype A Immunogen HIV-1 exposed seronegative

Species (MHC) human (B53)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix – systemic CD8+ T-cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T-cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T-cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

**HXB2 Location** p24 (47–58) Author Location p24 (181–192) Epitope CTPYDINQMLNC Immunogen HIV-2 infection Species (MHC) human (B58) References Bertoletti 1998

• HIV-2 epitope defined from an infection in Gambia, Bertoletti, pers. comm.

**HXB2 Location** p24 (47–58) **Author Location** p24 **Epitope** ATPQDLNTMLNT Subtype B, D

Immunogen HIV-1 infection Species (MHC) human (B58)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7 Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ Keywords subtype comparisons

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence in this person matched the known epitope.

**HXB2 Location** p24 (48–55)

**Author Location** p24 (48–55)

**Epitope** TPQDLNTM

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 7 patients recognized this epitope.

**HXB2 Location** p24 (48–56)

**Author Location** p24 (180–188 IIIB)

Epitope TPQDLNTML

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*0702 epitope.

**HXB2 Location** p24 (48–56)

Author Location (C consensus)

Epitope TPQDLNTML

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*3910)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the L9 residue of TPQDLNTML are associated with the presence of the HLA presenting molecule in the host.
- TPQDLNTML is cross-presented by B\*8101 and B\*3901.

**HXB2 Location** p24 (48–56)

**Author Location** 

**Epitope** TPQDLNTML

Immunogen

Species (MHC) human (B\*3910)

Keywords optimal epitope

**References** Frahm et al. 2007

• C. Brander notes that this is an B\*3910 epitope.

**HXB2 Location** p24 (48–56)

**Author Location** p24 (179–187 LAI)

Epitope TPQDLNTML

Subtype B

Immunogen

Species (MHC) human (B\*4201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*4201 epitope.

**HXB2 Location** p24 (48–56)

**Author Location** (C consensus)

Epitope TPQDLNTML

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- TPQDLNTML is an optimal epitope.

**HXB2 Location** p24 (48–56)

Author Location Gag (96ZM651.8)

Epitope TPQDLNTML

Epitope name G180-TL9

Immunogen

**Species (MHC)** human (B\*4201, B\*8101)

Keywords subtype comparisons, immunodominance

References Novitsky et al. 2001

 This study provides a survey of CTL responses and full length HIV-1 genome sequences from a C subtype infected Botswanan cohort.

- 19/46 (41.3%) had CTL responses to one or more peptides within the first immunodominant region of Gag (peptides TL-NAWVKVIEEKAFSPEVIP, EKAFSPEVIPMFTALSEGAT, and MFTALSEGATPQDLNTMLNT), with magnitudes of response with ELISPOT results median and range 495 (103 to 1,447) SFC/10<sup>6</sup> PBMC
- 7/11 HLA-A\*4201+ subjects (64%) responded to peptide MF-TALSEGATPODLNTMLNT.
- TPQDLNTML is a A\*4201 epitope within TL-NAWVKVIEEKAFSPEVIP.

**HXB2 Location** p24 (48-56)

Author Location p24

Epitope TPQDLNTML

Epitope name TL-9

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*4201, B\*8101)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Eli-

spot - IFNγ, Intracellular cytokine staining,

Chromium-release assay

Keywords subtype comparisons, epitope processing, im-

munodominance, cross-presentation by differ-

ent HLA

References Masemola et al. 2004b

- Highly targeted regions in Gag for CD8+ T-cells were defined for individuals with C clade infections in South Africa. 9 specific epitopes within the most reactive regions were characterized. Some epitopes were shown to be promiscuous, presented by multiple class I restricting alleles.
- TPQDLNTML was presented by B\*4201 and B\*8101. B\*44 is more common among Caucasians than Zulus (allele frequency 0.149 versus 0.107), while A\*29 is more common in Zulus (0.045 versus 0.125). This epitope had previously identified in B clade infections.

**HXB2 Location** p24 (48–56)

Author Location Gag (173-181 HIV-2)

Epitope TPYDINQML

Immunogen HIV-2 infection

Species (MHC) human (B\*5301)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*5301 epitope.

**HXB2 Location** p24 (48–56)

**Author Location** (48–56)

**Epitope** TPQDLNTML

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*81)

Assay type Other

Keywords epitope processing, HLA associated polymor-

phism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- TPQDLNTML was a previously defined B\*81 presented epitope that encompassed associated polymorphisms, SeGA|TPQDLNtML, in the seventh position of as well in the third position before the known epitope.

HXB2 Location p24 (48–56) Author Location p24 (180–188 LAI) Epitope TPQDLNTML Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*8101)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*8101 epitope.

HXB2 Location p24 (48–56) Author Location (C consensus) Epitope TPQDLNTML Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*8101) Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the T7 residue of TPQDLNTML are associated with the presence of the HLA presenting molecule in the host.
- TPQDLNTML is cross-presented by B\*8101 and B\*3901.

**HXB2 Location** p24 (48–56)

**Author Location** 

**Epitope** TPQDLNTML **Epitope name** Gag-TL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*8101, B\*5301, B7)

**Donor MHC** A\*3402 A\*7401 B\*5301 B\*8101 Cw\*0401 Cw\*0802

**Keywords** HAART, ART **References** Sabbaj *et al.* 2003

• This study monitored epitope responses in HIV-1 infected minority women living in the United States.

- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- Subjects 00RCH86 and 03RCH59 both recognized this epitope, both restricted by HLA B\*8101.
- Subject 00RCH86 was African American, not on HAART, viral load 51000, CD4 count 520.
- Subject 03RCH59 was African American, male, on HAART, viral load 22000, CD4 count 769.
- Among HIV+ individuals who carried HLA B07, 2/9 (22%) recognized this epitope.
- Among HIV+ individuals who carried HLA B\*5301, 3/15 (20%) recognized this epitope.
- Among HIV+ individuals who carried HLA B81, 4/6 (67%) recognized this epitope.

HXB2 Location p24 (48–56) Author Location (C consensus) Epitope TPQDLNTML

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B39, B\*4201, B\*8101, Cw\*0802)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** cross-presentation by different HLA, characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (48–56)

Author Location p24 (C consensus)

**Epitope** TPQDLNTML **Immunogen** HIV-1 infection

Species (MHC) human (B42)

**Keywords** subtype comparisons, immunodominance **References** Goulder *et al.* 2000a

B42 and B81 are very similar, and both can present this epitope to B42-positive effector cells – this epitope is almost certainly optimal for B81 as well – B42 and or B81 are expressed in 40-45% of Zulu and Xhosa infected individuals in South Africa, and in 14/18 B42 or B81+ individuals, the dominant gag response was to TPQDLNTML.

- SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31/44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEOA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32/37 C-clade infected subjects.

HXB2 Location p24 (48-56)

**Author Location** Gag

Epitope TPQDLNTML Immunogen HIV-1 infection Species (MHC) human (B42)

References Goulder et al. 2000b

- Tetramer assays were compared with three functional assays in 42 people with chronic HIV infection: ELISPOT, intracellular cytokine staining, and precursor frequency (limiting dilution assay [LDA]).
- HIV-specific tetramer staining CTLs appeared to be active, and inert CTL were not found to play a significant role in chronic pediatric or adult HIV infection.

**HXB2 Location** p24 (48–56)

Author Location p24

**Epitope** TPQDLNQML

**Immunogen** 

Species (MHC) human (B53)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied - these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive, and the B35 allele seems to be protective.
- HIV-2 sequence: TPYDINQML, no cross-reactivity, Gotch et al. [1993]

**HXB2 Location** p24 (48–56)

Author Location Gag (173–181 HIV-2)

Epitope TPYDINQML Immunogen HIV-2 infection Species (MHC) human (B53) References Gotch et al. 1993

**HXB2 Location** p24 (48–56)

Author Location Gag (180-188 subtype A)

Epitope TPQDLNMML

Subtype A

Immunogen HIV-1 infection, in vitro stimulation or selec-

tion

Species (MHC) human (B53)

**Keywords** subtype comparisons References Dorrell et al. 2001

• Three peptides GSEELRSLYNTVATL (p17 residues 71-85), • In vitro restimulation of CTL specific for dominant epitopes from infected individuals is possible using recombinant modified vaccinia virus Ankara (MVA) carrying A or D subtype HIV-1 Gag proteins.

**HXB2 Location** p24 (48–56)

Author Location p24 (180–188 subtype A consensus)

**Epitope** TPQDLNMML

Subtype A

Immunogen HIV-1 infection Species (MHC) human (B53)

> Keywords subtype comparisons, immunodominance, TCR usage

References Dorrell et al. 2001

- In clade A infected Gambians, three HLA-B53 epitopes were defined in Gag p24 using ELISPOT, tetramer, and cytotoxicity assays.
- This optimal epitope was identified within the 20 mer reactive peptide that carried it by homology with a B53 epitope from HIV-2, a B subtype B7 peptide that corresponds to it, as B53 is part of the B7 superfamily, and by the proline in the anchor at position 2.
- TPQDLNMML was recognized in 6/7 HLA-B53 subjects and was immunodominant in most subjects.
- TPQDLNMML was A subtype-specific with no crossrecognition of the subtype B, C, and D variant, TPQDLNTML, although the B/C/D variant bound more efficiently to B53 position 7 show great positional variation in crystal structures of two HLA-B53 complexes, suggesting variation here might significantly alter the position of the peptide in the binding groove and thus affect TCR interactions.
- Only one subject might have had a cross-reactive response with the HIV-2 and Mamu-A\*01 variant CTPYDINQML, and this subject might have been dual infected with HIV-2.

**HXB2** Location p24 (48–56)

**Author Location** p24

Epitope TPQDLNMML **Immunogen** HIV-1 infection Species (MHC) human (B53)

Assay type Intracellular cytokine staining

Keywords immunodominance, genital and mucosal immunity

References Kaul et al. 2003

- Predefined immunodominant peptide responses were used to compare CD8+ T cells responses in the blood and cervix of 16 HIV+ Kenyan sex workers. Cervical responses were detected in 8/10 women from whom adequate samples could be obtained. The frequency of the CD8+ T cell response in the genital tract was comparable to the blood, with a trend toward being slightly
- The immunodominant response was to this epitope in the PBMC of 2/16 patients (Kaul et al. 2001, AIDS, 107:1303).

**HXB2 Location** p24 (48-56)

Author Location Gag (180-188)

Epitope TPQDLNMML

Epitope name TPQ

Immunogen HIV-1 infection, HIV-2 infection

Species (MHC) human (B53)

Country Gambia.

Assay type CD8 T-cell Elispot - IFNγ, Tetramer binding, Flow cytometric T-cell cytokine assay

**Keywords** characterizing CD8+ T cells **References** Gillespie *et al.* 2005

- CD8 T-cell responses were evaluated and compared in HIV-1 and HIV-2 infected individuals living in the Gambia. A significantly greater magnitude and breadth of Gag-specific responses were found in HIV-1 infected individuals, possibly because of low or undetectable viral loads in HIV-2 infected individuals. This finding suggests that responses in HIV-2 infection reflect antigen load in plasma, as is the case in HIV-1 infection. No correlation was found between immune control of HIV-2 and the frequency of perforin-expressing virus-specific CD8 T-cells.
- 3/3 HIV-2-infected B53-positive subjects responded to the TPY-DINQML HIV-2 version of the epitope.
- 5/7 HIV-1-infected B53-positive subjects recognized both the HIV-1 equivalent epitope, TPQDLNMML, and also the QAS epitope.

**HXB2 Location** p24 (48–56)

Author Location Gag (180-188)

Epitope TPYDINQML

Epitope name TPY

Subtype HIV-2

Immunogen HIV-1 infection, HIV-2 infection

Species (MHC) human (B53)

Country Gambia.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Tetramer binding, Flow cytometric T-cell cytokine assay

**Keywords** characterizing CD8+ T cells

References Gillespie et al. 2005

- CD8 T-cell responses were evaluated and compared in HIV-1 and HIV-2 infected individuals living in the Gambia. A significantly greater magnitude and breadth of Gag-specific responses were found in HIV-1 infected individuals, possibly because of low or undetectable viral loads in HIV-2 infected individuals. This finding suggests that responses in HIV-2 infection reflect antigen load in plasma, as is the case in HIV-1 infection. No correlation was found between immune control of HIV-2 and the frequency of perforin-expressing virus-specific CD8 T-cells.
- 3/3 HIV-2-infected B53-positive subjects responded to the TPY-DINQML HIV-2 version of the epitope.
- 5/7 HIV-1-infected B53-positive subjects recognized both the HIV-1 equivalent epitope, TPQDLNMML, and also the QAS epitope.

**HXB2 Location** p24 (48–56)

Author Location p24 (180–188 IIIB)

Epitope TPQDLNTML

Immunogen HIV-1 infection Species (MHC) human (B7)

**Keywords** responses in children, mother-to-infant transmission, escape

References Wilson et al. 1999a

 This study describes maternal CTL responses in the context of mother-to-infant transmission.

- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- No variants of this epitope were found in a non-transmitting mother that had a CTL response to this epitope.

HXB2 Location p24 (48–56)

**Author Location** p24 (180–188)

Epitope TPQDLNTML

Immunogen HIV-1 infection

Species (MHC) human (B7)

 ${\bf Keywords} \ {\bf immunodominance}$ 

References Jin et al. 2000b

- This is the optimal epitope for the immunodominant response defined using a conventional approach in an HLA B7+ longterm non-progressor.
- Three additional sub-dominant HLA B7 epitopes were defined using EpiMatrix, a non-anchor based strategy for defining potential epitopes, which highlighted 2078 possible epitopes in the autologous HIV-1 derived from the study subject – this was followed by B7 anchor residue prediction which narrowed the set to 55 peptides, three of which could serve as functional CTL epitopes.

**HXB2 Location** p24 (48–56)

**Author Location** p24 (SF2)

Epitope TPQDLNTML

Epitope name TL9

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Goulder et al. 2001a

 Recognized by patient 9354 during chronic infection, used as a positive control in a study of the SLYNTVATL epitope.

**HXB2 Location** p24 (48–56)

**Author Location** p24 (48–56)

Epitope TPQDLNTML

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

**HXB2 Location** p24 (48–56)

Author Location p24 (48–56)
Epitope TPQDLNTML
Epitope name B7-TL9
Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7) Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 1/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 1/4 individuals had detectable responses to this epitope after STI.

**HXB2 Location** p24 (48–56)

Author Location p24

Epitope TPQDLNTML
Epitope name B7-TL9(p24)

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7) Donor MHC A32, B7, B14

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8+ T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals. Patient A displayed the greatest response to epitope B14-EL9(gp41), a strong response to B7-TL9(p24), and responses to B7-TM9(Nef) and A32-PW10(RT).

**HXB2 Location** p24 (48–56)

Author Location p24 (48-56)

Epitope TPQDLNTML

**Epitope name** B7-TL9 Gag

**Subtype** B

Immunogen HIV-1 infection Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

 An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response, but in this case the epitope did not vary.

**HXB2 Location** p24 (48–56)

Author Location (B consensus)

Epitope TPQDLNTML

Epitope name TL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A03, B07, Cw7

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope

**HXB2 Location** p24 (48–56)

Author Location p24 (HXB2)

Epitope TPQDLNTML

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (B7, Cw8, B42)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** epitope processing, vaccine antigen design, characterizing CD8+ T cells

References SenGupta et al. 2004

Multiple HLA class I-restricted and class II-restricted T-cell
epitopes were shown to be processed and presented from an
exogenously added HIV-1 gag-p24 peptide complexed to a
heat shock protein. T-cell recognition of the complex was
shown to be inhibited by brefeldin A indicating an endoplasmic
reticulum-dependent pathway.

**HXB2 Location** p24 (48–56)

Author Location p24 (180–188 LAI)

Epitope TPQDLNTML

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0802)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a C\*0802(Cw8) epitope.

**HXB2 Location** p24 (48–56)

**Author Location** (C consensus)

**Epitope** TPQDLNTML

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0802)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- TPQDLNTML is an optimal epitope.

**HXB2 Location** p24 (48–56)

Author Location Gag (180-188)

**Epitope** TPQDLNTML

Epitope name TL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw8)

**Donor MHC** A\*02, A\*68, B\*14, B\*52, Cw\*08, Cw\*12

**Country** United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords escape, optimal epitope

References Koibuchi et al. 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- The B consensus form of this epitope, TPQDLNTML, persisted throughout 6 years of chronic infection in 1 individual.

**HXB2 Location** p24 (48–56)

**Author Location** Gag

Epitope TPQDLNTML

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFNγ

Keywords assay standardization/improvement, epitope

processing, characterizing CD8+ T cells

References Beattie et al. 2004

- This study compared CD8+ T-cell EliSpot reponses to 58 Gag peptides that were optimal epitopes, with responses to overlapping 15 mers that spanned Gag. When screening for HIV-1-specific CD8+ T-cell responses from 49 HIV+ people, overlapping 15-mer peptide pools revealed several novel responses that would have been missed using predefined CD8 epitopes. However, the 15-mer pools often missed low-level responses to predefined epitopes, especially when the epitope was located centrally in the 15-mer peptide, and the overall level of response to the 15 mers was generally lower(mean 1.4 five fold dilutions lower, range 0-3).
- The response to TPQDLNTML was used as an example of a titration curve. When comparing the peptide TPQDLNTML to the 15 mer EGATPQDLNTMLNTV, the 15 mer had a diminished response to the same amount of peptide.

**HXB2 Location** p24 (48–57)

**Author Location** Gag

Epitope TPQDLNMMLN

Immunogen

Species (MHC) human (B7)

References De Groot et al. 2001

- The program Epimatrix was used in conjunction with the program Conservatrix to identify conserved regions of HIV that might serve as epitopes.
- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes were identified that could stimulate IFNγ production in an ELISPOT assay.
- TPQDLNMMLN was newly defined as an HLA-B7 epitope in this study, athough it was previously published as a B\*8101 epitope.
- TPQDLNMMLN was shown to stimulate an ELISPOT response, but could not be shown to bind to HLA-B7.
- The variant TPQDLNTMLN was cross-reactive, had previously been identified as a HLA-B14 epitope, and could bind to HLA-B7.

**HXB2 Location** p24 (48–57)

**Author Location** Gag

**Epitope** TPQDLNMMLN

Epitope name 1309

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A03, A24, B07, B38, Cw07, Cw12/13

Country United States.

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope prediction

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for TPQDLNMMLN: 31%.

**HXB2 Location** p24 (48–57)

**Author Location** Gag

Epitope TPQDLNTMLN

Epitope name 1308

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (B7, B14)

Country United States.

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope prediction, cross-presentation by different HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for TPQDLNTMLN: 31%. This epitope was not confirmed in this study, but has been reported to be a B14 epitope.

**HXB2 Location** p24 (48-59)

Author Location p24

**Epitope** TPQDLNQMLNTV

Subtype B. G

Immunogen HIV-1 infection

Species (MHC) human (B58)

Donor MHC A2, A36, B45, B58, Cw3, Cw6

Country Nigeria.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, variant cross recognition or cross-neutralization

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype G Gag. The autologous epitope sequence in this person differed from the known epitope in one position, Q7T, TPQDLNtMLNTV

**HXB2 Location** p24 (48–62)

Author Location p24 (48-56)

Epitope TPQDLNMMLNIVGGH

Subtype A, D

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*0201, A\*2902, B\*1402, B\*1503; A\*0101,

A\*7401, B\*5801

Country Uganda.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, variant cross-

recognition or cross-neutralization

References Barugahare et al. 2005

- T-cell immune responses were examined in HIV-1 infected Ugandans. Similar levels of cross-clade responses for Gag, Env and Nef were observed. Higher frequencies of responses in conserved regions were found for Gag within a single clade, while areas with higher viral sequence variation had fewer detectable responses. Differential interclade sequence homologies between the Gag regions did not affect the respective level of cross-clade recognition.
- This sequence contains a known B7 and B53 epitope, but the subjects recognizing it are B7- and B53-negative. It was conserved in the two people that recognized the peptide.

**HXB2 Location** p24 (49–57)

Author Location p24 (181–189 LAI)

**Epitope** PQDLNTMLN

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14, Cw8)

References Lubaki et al. 1997

- Eighty two HIV-1-specific CTL clones from 5 long-term nonprogressors were isolated and analyzed for breadth of response.
- A sustained Gag, Env and Nef response was observed, and clones were restricted by multiple HLA epitopes, indicating a polyclonal response.
- Despite this being a well defined conserved epitope, none of the 11 gag-specific clones from a B-14 positive subject could recognize either it or p24 RAEQASQEV.
- Christian Brander notes that B14 and Cw8 are in linkage disequilibrium, and that this epitope may be Cw8.

HXB2 Location p24 (51-59)

**Author Location** p24 (subtype A)

Epitope DLNMMLNIV

Subtype A

**Immunogen** HIV-1 exposed seronegative

Species (MHC) human (B14)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix systemic CD8+ T-cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T-cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T-cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

**HXB2 Location** p24 (51–59)

**Author Location** p24

Epitope DLNMMLNIV

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Keywords** HIV exposed persistently seronegative (HEPS)

## References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted - 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls, ML1792.

**HXB2 Location** p24 (51–59)

Author Location p24 (183–191 LAI)

**Epitope** DLNTMLNTV

Epitope name G5

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords HAART, ART

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened - eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+ PBL - but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

HXB2 Location p24 (51-59)

**Author Location** p24 (183–191)

Epitope DLNMMLNIV

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B14)

Keywords HIV exposed persistently seronegative

(HEPS), immunodominance

References Kaul et al. 2001a

- Variants DLNMMLNIV/DLNTMLNVV are specific for clades
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.

- Among HLA-B14 women, 4/4 HEPS and 3/7 HIV-1 infected women recognized this epitope, likelihood ratio 4.8, p value 0.1, and HEPS women tended to respond to DLNMML-NIV/DLNTMLNVV, while infected women tended to respond to DRF(F/W)KTLRA.
- The dominant response to this HLA allele was to this epitope for all 4/4 HEPS cases and in only one of the 3/7 HIV-1 infected
- Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.
- Four epitopes were considered to be "resistant epitopes", as they were preferentially reactive in HEPS women and so may confer resistance, and these were found in three different proteins: A2 ILK(D/E)PVHGV in RT, A\*6802 DTVLEDINL in Protease, B14 DLN(M/T)LN(I/V)V in p24 and B18 FRDYV-DRF(Y/F)K also in p24.

**HXB2 Location** p24 (51–59)

Author Location p24

**Epitope** DLNMMLNIV

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

**HXB2 Location** p24 (51–59)

Author Location p24 (183-191 LAI)

**Epitope** DLNTMLNTV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14, Cw8)

References Johnson et al. 1992; Nixon et al. 1988

Recent evidence indicates this is a Cw8 epitope; B14 and Cw8 are in linkage disequilibrium and the HLA presenting molecule is hard to distinguish (P. Goulder, personal communication).

**HXB2 Location** p24 (51–59)

Author Location p24

**Epitope** DLNTMLNTV

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B14, Cw8)

Keywords subtype comparisons, HIV exposed persis-

tently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades such cross-reactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A subtype consensus is identical to the B clade epitope.
- The D subtype consensus is dLNmMLNiV.
- Recent evidence indicates this is a Cw8 epitope;B14 and Cw8
  are in linkage disequilibrium and the HLA presenting molecule
  is hard to distinguish (P. Goulder, personal communication).

**HXB2 Location** p24 (51–59)

Author Location p24 (183-191 LAI)

**Epitope** DLNTMLNTV

Subtype B

Immunogen HIV-1 infection Species (MHC) human (Cw8)

Keywords review

References McMichael & Walker 1994

- Review of HIV CTL epitopes defined by B14 motif found within a larger peptide.
- Recent evidence indicates this is a Cw8 epitope;B14 and Cw8 are in linkage disequilibrium and the HLA presenting molecule is hard to distinguish (P. Goulder, personal communication).

**HXB2 Location** p24 (51–59)

Author Location p24 (subtype B)

**Epitope** DLNTMLNTV

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (Cw8, B\*1402)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among B and D clade viruses.
- The Clade A version of the epitope, DLNNMLNIV, was preferentially recognized by CTL.
- Recent evidence indicates this is a Cw8 epitope; B14 and Cw8 are in linkage disequilibrium and the HLA presenting molecule is hard to distinguish (P. Goulder, personal communication).

**HXB2 Location** p24 (51–59)

Author Location p24

Epitope DLNTMLNTV Immunogen HIV-1 infection Species (MHC) chimpanzee References Santra *et al.* 1999

- 3/4 animals displayed HIV-1 Gag-specific CTL activity.
- Effector cells from two chimpanzees were able to recognize two epitopes also recognized by human HIV-1 Gag-specific CTL (SPRTLNAWV, HLA-B7, and DLNTMLNTV, HLA-B14).

 No chimpanzee CTL were detected to the following human HIV-1 specific Gag epitopes, although they were embedded within 20mer peptides that contained a reactive epitope: ISPRTLNAW, HLA-B57; KRWIILGLNK, HLA-B27; and DRFYKTLRA, HLA-B14.

HXB2 Location p24 (51-60)

**Author Location** Gag

Epitope DLNTMLNTVG

Epitope name 1238 Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A2, B14)

**Country** United States. **Assay type** T-cell Elispot

**Keywords** binding affinity, computational epitope prediction, cross-presentation by different HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for DLNTMLNTVG: 65%. This
  epitope was not confirmed in this study, but was previously
  reported to be presented by B14.

**HXB2 Location** p24 (51–70)

Author Location p24 (183-202 SF2)

Epitope DLNTMLNTVGGHQAAMQMLK

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A26, A30, B38.

**HXB2 Location** p24 (51–82)

**Author Location** Gag (183–214 LAI)

Epitope DLNTMLNTVGGHQAAMQMLKETINEEAAEWDR

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long amino acid peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I trial.
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees – 2/10 reacted to this peptide.
- 9/12 tested mounted a CTL response to at least one of the six peptides; each of the six peptides elicited a CTL response in at least one individual.

• None of the 12 tested had an IgG response to this peptide.

Author Location p24 (61–69)
Author Location p24 (61–69)
Epitope GHQAAMQML
Immunogen HIV-1 infection
Species (MHC) human (B\*1510)
Keywords optimal epitope
References Frahm et al. 2007

HXB2 Location p24 (61–69)
Author Location (C consensus)
Epitope GHQAAMQML
Subtype C
Immunogen HIV-1 infection
Species (MHC) human (B\*1510)
Country South Africa.

**Assay type** CD8 T-cell Elispot - IFNγ **Keywords** characterizing CD8+ T cells **References** Kiepiela *et al.* 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location p24 (61–69)

Author Location (C consensus)

Epitope GHQAAMQML

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1510)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

 A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses

were associated with higher viremia.

• GHQAAMQML is an optimal epitope.

HXB2 Location p24 (61–69)
Author Location p24 (193–201 LAI)
Epitope GHQAAMQML
Subtype B
Immunogen
Species (MHC) human (B\*3901)

**References** Frahm *et al.* 2007
• C. Brander notes this is a B\*3901 epitope.

Keywords optimal epitope

HXB2 Location p24 (61–69)
Author Location Gag (193–201 IIIB)
Epitope GHQAAMQML
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (B38)
Assay type Chromium-release assay
References Kurane et al. 2003

Three CD8+ CTL cell clones were derived from 2 HIV-1 positive asymptomatic patients, and their epitope specificities and HLA presenting proteins were defined.

HXB2 Location p24 (61–69)
Author Location p24 (193–201 LAI)
Epitope GHQAAMQML
Subtype B
Immunogen
Species (MHC) human (B39)
References Kurane & West 1998

• Optimal peptide defined by titration.

HXB2 Location p24 (61–71)

Author Location p24 (193–203 BRU)

Epitope GHQAAMQMLKE

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Claverie et al. 1988

 One of 4 epitopes first predicted, then shown to stimulate HLA-A2 restricted CTL line.

HXB2 Location p24 (61–71)
Author Location p24 (61–70)
Epitope GHQAAMQMLKE
Immunogen HIV-1 infection
Species (MHC) human (A2)
Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses.
   HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 7/19 patients recognized this epitope.

HXB2 Location p24 (61–80)
Author Location p24 (193–212 SF2)
Epitope GHQAAMQMKETINEEAAEW
Immunogen HIV-1 infection
Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.

• The responding subject was HLA-A26, A30, B38.

HXB2 Location p24 (61-82)

**Author Location** p24 (193-214 BH10)

Epitope GHQAAMQMLKETINEEAAEWDR

Immunogen HIV-1 infection Species (MHC) human (B52) References Johnson *et al.* 1991

• Gag CTL response studied in three individuals.

**HXB2 Location** p24 (62–70)

Author Location p24 (194-202 LAI)

**Epitope** HQAAMQMLK

Subtype B

Immunogen

Species (MHC) human (B52)

**References** Brander & Walker 1996

• P. Goulder, pers. comm.

HXB2 Location p24 (64-80)

**Author Location** p24 (63–80 HXB2)

Epitope AAMQMLKETINEEAAEW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 22% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p24 (65–72)

Author Location p24

Epitope AMQMLKET

Epitope name A9I

Immunogen vaccine

Vector/Type: DNA HIV component: Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type Chromium-release assay

References Bojak et al. 2002b

 Codon-optimized gag gene DNA vaccines were compared to wild type by vaccination of BALB/c mice. Codon optimized DNA gave a Th1 polarized Th response, a strong antibody response that persisted from more than 20 weeks, and CTL responses, while wild-type DNA induced weak and inconsistent immune responses.

**HXB2 Location** p24 (65–73)

Author Location Gag (197-205 BRU)

Epitope AMQMLKETI Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 1/9 CRF02\_AG-infected Ivorians, and 1/9 B-infected French subjects.

**HXB2 Location** p24 (65–73)

Author Location Gag (199–207 HXB2)

Epitope AMQMLKETI

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade HXB2

HIV component: Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

**References** Qiu et al. 1999

- Different expression vectors were tested to increase Gag expression in cell lines and create suitable vectors for DNA vaccines.
- Stable Gag expression was achieved in murine p815 cells, using a Gag gene that had mutated silent base positions that disrupt inhibitory RNA sequences which promote RNA degradation.
- Silent mutations were more effective than introduction of the D retrovirus cis-acting posttranscriptional control element (CTE) for enhancing Gag expression.
- The gag vector with silent mutations given as a vaccine to BALB/c mice gave CTL responses in splenic mononuclear cells, using peptide pulsed cells as targets.

**HXB2 Location** p24 (65–73)

Author Location p24 (199–207 SF2)

Epitope AMQMLKETI

Epitope name p7g

Immunogen vaccine

Vector/Type: protein, vaccinia Strain: B clade SF2 HIV component: Gag, Gag-Pol Adjuvant: E. coli mutant heat labile entero-

toxin (LT-R72)

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Neidleman et al. 2000

- Intranasal immunization of CB6F1 (H2bxd) mice with soluble gag p55 with LT ADP-ribosyltransferase mutants (LTK63 and LTK73) from Escherichia coli as adjuvants was tested.
- Intranasal and intramucosal immunization of p55 gag protein with LTK63 or LTK72 adjuvant induced a CTL response comparable to intramuscular immunization responses.
- Oral co-administration of LTR72, with residual ADPribosyltransferase activity, induced systemic CTL responses, but LTK63 with no ADP-ribosyltransferase activity did not.

HXB2 Location p24 (65–73) Author Location p24 (66–74) Epitope AMQMLKETI Immunogen vaccine

Vector/Type: DNA HIV component: Gag Adjuvant: vesicular stomatitis virus glycoprotein (VSV-G)

Species (MHC) mouse (H-2<sup>d</sup>)
References Marsac *et al.* 2002

- BALB/c mice were injected with plasmids expressing HIV-1
  Gag with or without coinjection of a plasmid expressing vesicular stomatitis virus glycoprotein (VSV-G). The combination encodes VSV-G pseudotyped Gag particles that can be taken up by cells for presentation in either the class I or class II pathways, while exogenous Gag alone can only be taken into the class II pathway.
- Vaccination with DNA expressing VSV-G pseudotyped Gag particles rather than just Gag increase Gag-specific CTL responses generally as well as the specific H-2d restricted anti-AMQMLKETI response.

**HXB2 Location** p24 (65–73)

**Author Location** Gag

Epitope ANQMLKDTI

Subtype C

Immunogen vaccine

*Vector/Type:* DNA with CMV promotor *HIV component:* Gag, Protease

**Species (MHC)** mouse (H-2<sup>d</sup>)

Country India.

Assay type Cytokine production, CD8 T-cell Elispot -

IFN $\gamma$ , CD4 T-cell Elispot - IFN $\gamma$ , Cytolytic LDH release assay

**Keywords** subtype comparisons, vaccine-induced epitopes, Th1

References Chugh & Seth 2004

- A gag-protease gene construct from HIV-1 subtype C Indian strain has been shown to be successful in evoking immune responses to gag epitopes from both CD4+ and CD8+ T-cells in BALB/c mice. The immune response was of TH1 type. Recognition of seven Gag peptides carrying multiple epitopes indicates a broad-based immune response.
- A cross-clade response to the C clade epitope ANQMLKDTI was observed to the B clade version of this epitope, aNqmlkEti.
   66% lysis was observed to the peptide carrying the C clade epitope, only 33% to the B clade variant.

HXB2 Location p24 (65–73) Author Location p24 (199–207 SF2) Epitope AMQMLKETI Epitope name p7G Immunogen vaccine

Vector/Type: protein Strain: B clade SF2 HIV component: Gag Adjuvant: DDA, DOTAP, CpG immunostimulatory sequence (ISS), MF59, PLG, urea

Species (MHC) mouse (H-2<sup>kd</sup>)

Keywords dendritic cells

References O'Hagan et al. 2002

- Intramuscular or intraperitoneal immunization of BALB/c or CB6F1 mice with urea-solubilized, emulsified, or PLGmicroparticle associated p55 Gag was studied in conjunction with the adjuvant CpG. CpG did not enhance CTL immunity when combined with urea solubilized p55, but did when combined with emulsions and PLG-microparticle antigen.
- CpG shifted the Ab response towards a IgG2a, and CpG was shown to upregulate CD86 on mouse bone-marrow derived dendritic cells.

**HXB2 Location** p24 (65–73)

**Author Location** p24 (199–207 SF2)

Epitope AMQMLKETI

Subtype B

Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade SF2 HIV component: Gag, gp120

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Assay type Chromium-release assay

Keywords epitope processing, vaccine-induced epitopes

References Doe et al. 1996

 Spleen cells from mice with distinct MHC types were infused into HIV vaccinated scid mice, to study the antigen presenting cells used by CTL induced in intramuscular injections. Bone marrow derived cells are used for presenation, but DNA infection is not required for priming, rather APCs can present proteins synthesized in other host cells.

**HXB2 Location** p24 (65–73)

Author Location p24 (199–207 SF2)

Epitope AMQMLKETI

Immunogen vaccine

Vector/Type: vaccinia HIV component: Gag, Pol

**Species (MHC)** mouse (H-2K<sup>d</sup>)

Keywords immunodominance

**References** Doe & Walker 1996

- Immunodominant murine CTL response to this peptide observed after immunization with vaccine VVgagpol.
- Optimal peptide was defined.

**HXB2 Location** p24 (65–73)

**Author Location** Gag (197–205)

Epitope AMQMLKETI

Immunogen vaccine

Vector/Type: Listeria monocytogenes HIV

component: Gag

**Species (MHC)** mouse (H-2K<sup>d</sup>)

References Rayevskaya & Frankel 2001

- binant Listeria monocytogenes, Lmdaldat, that can grow only when supplemented with D-alanine, and that expresses HIV-1 HXB2 Gag.
- Parenteral immunization provided protection against systemic and mucosal challenges with a recombinant vaccinia virus expressing HIV-1 gag, and a long lasting memory CTL response against Gag in spleen, mesenteric lymph nodes, and Pever's patches directed against the gag protein.
- Oral immunization gave protection only against mucosal virus challenge and was associated with a transient CTL response in the three lymphoid tissues examined.
- L. monocytogenes is a gram-positive bacteria that enters the macrophage on phagocytosis and lives in the cytoplasm – secreted L. monocytogenes antigens are processed and presented by both class I and class II pathways.

HXB2 Location p24 (65-73)

**Author Location** Gag (197–205 SF2)

Epitope AMQMLKETI Immunogen vaccine

> Vector/Type: Listeria monocytogenes Strain: B clade HXB2 HIV component: Gag

**Species (MHC)** mouse (H-2K<sup>d</sup>)

Keywords immunodominance

References Mata et al. 1998

- BALB/c mice were immunized with recombinant Listeria monocytogenes (Lm-Gag) expressing HIV-1 HXB2 Gag.
- L. monocytogenes is a gram-positive bacteria that enters the macrophage on phagocytosis and lives in the cytoplasm - secreted L. monocytogenes antigens are processed and presented by both class I and class II pathways.
- This is the immunodominant CTL epitope in Gag in BALB/c mice.
- AMQMLKETI does not contain established Kd anchoring residue in position 2, tyrosine or phenylalanine, thus deviating from the typical Kd anchoring motif - the lack of the aromatic anchor residue is compensated for by interaction of the glutamine at P3 with pocket D of Kd.

**HXB2 Location** p24 (65–73)

Author Location Gag (HXB2)

**Epitope** AMQMLKETI

Subtype B

Immunogen vaccine

Vector/Type: vaccinia, vesicular stomatitis virus (VSV) Strain: B clade HXB2, B clade

IIIB HIV component: Env, Gag

**Species (MHC)** mouse (H-2K<sup>d</sup>)

Keywords immunodominance References Haglund et al. 2002a

- Different HIV strains were used for different regions: Env IIIB, Gag HXB2.
- BALB/c mice were vaccinated with rec vesicular stomatitis virus (rVSV) expressing either HIV-1 Gag, Env, or both, and compared to using rec Env and Gag in vaccinia virus (rVVs). The primary response was determined by cell lysis, cytokine production and tetramer staining.

- BALB/c mice were immunized with a highly attenuated recom• Primary CTL responses to the immunodominant Gag (AMOM-LKETI) epitope peaked in 7 days for GAG-rVSV, 3% of the cells were tetramer positive, and this response was 8-fold higher than for Gag-rVV.
  - Vaccinating with GagEnv-rVSV carrying both Gag and Env allowed recognition of both HIV-1 proteins, but at reduced levels compared to either Gag-rVSV or Env-rVSV alone.
  - Intranasal immunization with Env-rVSV yielded CTL responses that were strong but reduced compared to an intraperitoneal route.

**HXB2 Location** p24 (65–73)

Author Location Gag (HXB2)

Epitope AMQMLKETI

Subtype B

Immunogen vaccine

Vector/Type: vaccinia, vesicular stomatitis virus (VSV) Strain: B clade HXB2, B clade

IIIB HIV component: Env, Gag

**Species (MHC)** mouse (H-2K<sup>d</sup>)

Keywords immunodominance

References Haglund et al. 2002b

- Different HIV strains were used for different regions: Env IIIB, Gag HXB2.
- BALB/c mice were vaccinated with rec vesicular stomatitis virus (rVSV) expressing either HIV-1 Gag or Env, or both, and retention of memory responses and recall responses were studied by tetramer staining and IFN-gamma production.
- · Seven months after vaccination with Env-rVSV, 6% of the CD8+ cells were tetramer positive for the immunodominant Env epitope; these cells had a memory phenotype, CD44-Hi
- Env in rec vaccinia virus (Env-rVV) elicited a strong recall response, with up to 45% to the CD8+ T-cell population tetramer positive and activated (expressing CD62L-Lo), and capable of IFN-gamma production.
- A prime with Env-rVSV and heterologous boost of Env-rVV gave remarkably high levels of memory cells, with approximately 1/3 of the CD8+ splenocytes being Env specific memory cells 150 days after the boost.
- A Gag-rVSV or EnvGag-rVSV prime and with a heterologous Gag-rVV or EnvGag-rVV boost combination gave 40% tetramer positive CD8+ cells, but the fraction of IFN-gamma producing cells was only about 25%. Still the heterologous vector prime-boost combination showed a profound benefit.
- A HIV-1 protein rVSV prime, rVV boost was a more potent combination than a vector reversal of a rVV prime and rVSV boost.

**HXB2 Location** p24 (65–73)

**Author Location** Gag

Epitope AMQMLKETI

Subtype B

Immunogen vaccine

Vector/Type: Listeria monocytogenes HIV component: Gag

**Species (MHC)** mouse (H-2K<sup>d</sup>)

Donor MHC H-2d

Assay type Tetramer binding, Intracellular cytokine stain-

**Keywords** genital and mucosal immunity **References** Peters *et al.* 2003

- Intravenous, rectal, and oral vaccination of recombinant L.
   monocytogenes expressing HIV-1 Gag antigen were compared for their ability to stimulate a mucosal CTL response; mucosal administration of this vaccine gave strong mucosal response that was readily boosted.
- This CTL epitope is the immunodominant epitope in Gag for BALB/c mice, and was used to characterize the vaccine responses.

**HXB2 Location** p24 (65–73)

Author Location Gag (197–205)

Epitope AMQMLKETI

Subtype B

Immunogen vaccine

Vector/Type: vaccinia, Listeria monocyto-

genes HIV component: Gag, Nef

**Species (MHC)** mouse (H-2K<sup>d</sup>)

Donor MHC H-2d

**Assay type** Cytokine production, proliferation, Tetramer binding, Intracellular cytokine staining,

Chromium-release assay

Keywords memory cells

References Rayevskaya et al. 2003

• Splenocytes derived from BALB/c mice immunized and boosted with Lmdd-gag were stimulated with gag-peptide specific antigen in vitro. In culture, CTL activity against this epitope reached a maximum at 9 days, then declined. Peptide restimulation gave a delayed (18 hours) but potent response, and growth was IL-2 or IL-15 dependent. Adoptive transfer of 5000 of the sorting purified cells could protect recipient BALB/c against vaccinia-gag challenge up to 3 months after transfer.

**HXB2 Location** p24 (65–73)

**Author Location** 

Epitope AMQMLKETI

Epitope name A9I Immunogen vaccine

Vector/Type: DNA, virus-like particle (VLP), polyepitope HIV component: Gag, p24 Gag,

V3

**Species (MHC)** mouse (H-2K<sup>d</sup>)

Assay type Cytokine production, Chromium-release as-

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance, vaccine

antigen design

References Wild et al. 2004

A codon optimized gag DNA vaccine was compared to a
myristylation defective gag and p24 alone, both of which lack
signals for secretion from transfected cells. Gag-derived immunogens that were secreted as VLPs and those that remained
intracellular (p24) each produced strong CTL responses, and
neither the size of antigen nor cellular trafficking and localization significantly influenced the strength of humoral and
cellular immune activation. The formation and release of VLPs
was not essential for eliciting strong CTL. BALB/c mice were

given the DNA vaccine by i.m. administration of plasmid DNA for the prime and boost.

- Linking the region encoding the V3 immunodominant epitope to the gag gene did not diminish the response to the Gag p24 epitope A9I, but did enable a response to the V3 epitope.
- Minigenes were made incorporating just 1 epitope, minitopes, carrying 1 of 3 murine class I eptiopes linked to the Ad2-E3 protein-derived signal peptide to allow access of the epitope to the ER. Weak induction of cellular immune responses was observed, in contrast to the complex polyprotein.

**HXB2 Location** p24 (65–73)

Author Location Gag (197-205)

Epitope AMQMLKETI

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade HXB2 HIV component: Gag

**Species (MHC)** mouse (H-2K<sup>d</sup>)

**Country** United States.

Assay type proliferation, T-cell Elispot

Keywords vaccine antigen design

References Kwak et al. 2004

 A recombinant vaccinia virus with HIV-1 Gag replacing the cytoplasmic domain of the B5R protein was shown to induce better primary CD4 response than recombinant vaccinia virus expressing Gag from the TK-locus; CD8 responses were less specific. When immunized BALB/c mice were challenged with a recombinant Listeria that expresses HIV-Gag, lower colony counts of Listeria were found in the liver and spleen of mice immunized with virus expressing B5R-Gag fusion protein.

**HXB2 Location** p24 (65–73)

**Author Location** Gag (199–207)

Epitope AMQMLKETI

Epitope name p7g

Subtype B

Immunogen vaccine

Vector/Type: vaccinia, Sindbis HIV compo-

nent: Gag

Species (MHC) mouse

**Assay type** CD8 T-cell Elispot - IFNγ

Keywords genital and mucosal immunity

References Vajdy et al. 2001

- Nasal, vaginal, rectal and i.m. immunization was performed with Sindbis virus expressing HIV-1 Gag (SIN-Gag), followed by intravaginal or intrarectal challenge with vaccinia virus expressing either Gag (VV-Gag) or gp160 (VV-gp160) as a control.
- Intranasal and intramuscular immunization followed by intravaginal challenge induced HIV-1 Gag specific, IFN-γ producing CD8+ T-cells in the vaginal/uteral mucosal tissue, as well as in the draining iliac lymph nodes and in the spleen, but could not protect against a VV-Gag infection of the ovaries. Local vaginal or rectal immunization, despite lower CD8+ T-cell responses, did provide protection.

**HXB2 Location** p24 (65–73)

**Author Location** Gag (Du422)

Epitope AMQMLKDTI

Subtype C

Immunogen vaccine

Vector/Type: DNA Strain: C clade Du422

HIV component: Gag

**Species (MHC)** mouse **Donor MHC** H-2d

Assay type Chromium-release assay

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References van Harmelen et al. 2003

- The pTHgagC DNA vaccine employed in this study expressed the gag gene derived from the South African isolate Du422, which was selected on the basis of being the natural strain most similar to the South African subtype C consensus sequence (aa distance of 1.8%).
- A E7D mutation was introduced into the epitope to match the gag subtype C sequence in the vaccine. Mice vaccinated with the gag DNA made strong CTL responses against AMQM-LKDTI, boosting enhanced the response, and memory cells persisted for 15 weeks.

**HXB2 Location** p24 (65–73)

Author Location p24 (197–205)

Epitope AMQMLKETI?

Immunogen vaccine

Vector/Type: protein HIV component: Gag

Adjuvant: Cholera toxin (CT)

Species (MHC) mouse

Donor MHC H-2d

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, Chromium-release assay

**Keywords** TCR usage, genital and mucosal immunity

References Yoshizawa et al. 2003

- Intranasal immunization triggered CTL response in the nasalassociated lymphoid tissue (NALT), posterior cervical lymph nodes (pCLNs) and the spleen, but not in the mesenteric lymph nodes (MLNs). Rectal immunzation elicited CTL responses only in the MLNs. By immunizing mice nasally following rectal immunization, CTL responses were detected in NALT, pCLNs, spleen and MLNs. Epitope-specific CD8+ T-cells were primarily located in NALT after 6 days and in pCLNs after 2 months.
- The strongest specific lysis was induced by NALT-specific CTL clones. pCLNs derived memory CTL clones originated from NALT CTL clones, as determined by T-cell receptor  $V\beta$  usage.

**HXB2 Location** p24 (69–86)

**Author Location** (C consensus)

Epitope LKDTINEEAAEWDRLHPV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** p24 (69–86)

**Author Location** Gag (201–218 LAI)

Epitope LKETINEEAAEWDRVPV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Buseyne et al. 1993a

- Vertical transmission of HIV ranges from 13% to 39%.
- Primary assays showed that cytotoxic activity against at least one HIV protein detected in 70% of infected children.
- Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.
- Patient EM28 (CDC P2A) had a CTL response to four epitopes in Gag.

**HXB2 Location** p24 (70–78)

**Author Location** 

Epitope KETINEEAA

Epitope name Gag-KA9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*4002)

 $\textbf{Donor MHC} \ \ 01RCH46: \quad \ A*0201, \quad A*0217, \quad B*0801,$ 

B\*4002, Cw\*0303, Cw\*0701

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- This epitope was newly defined in this study.
- Patient 01RCH46 was Hispanic, on HAART, and had a viral load of 21000 and CD4 count of 623 – she also recognized GELDRWEKI, p17(11-19), HLA B\*4002, and TAFTIPSI, RT(128-135), HLA A\*0217.
- Among HIV+ individuals who carried HLA B40, 3/5 (60%) recognized this epitope.

**HXB2 Location** p24 (70–78)

**Author Location** p24 (70–78)

Epitope KETINEEAA

Immunogen HIV-1 infection

Species (MHC) human (B\*4002)

Keywords optimal epitope

**References** Frahm *et al.* 2007

**HXB2 Location** p24 (71–80)

**Author Location** p24 (203–212)

Epitope ETINEEAAEW

Immunogen HIV-1 infection

Species (MHC) human (A\*2501)

**Keywords** subtype comparisons

**References** Klenerman et al. 1996

• The epitope was defined through direct stimulation of PBMC with 20-mer peptides.

- D. and E subtype isolates.
- DTINEEAAEW is found in A and some D subtype sequences.

HXB2 Location p24 (71-80) **Author Location** p24 (203–212) Epitope ETINEEAAEW Immunogen HIV-1 infection Species (MHC) human (A\*2501) Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is an A\*2501 epitope.

**HXB2 Location** p24 (71–80) **Author Location** p24 (203–212) **Epitope** ETINEEAAEW Immunogen HIV-1 infection Species (MHC) human (A\*2501) References van Baalen et al. 1996

- Conserved between B and D subtypes, variable in other clades; a consensus of clades A, C, F, G, and H and a peptide of HIV-2ROD over this region were not recognized by CTL recognizing the index peptide.
- C. Brander notes that this is an A\*2501 epitope in the 1999 database.

**HXB2 Location** p24 (71–80) Author Location p24 (71–80 HXB2) Epitope ETINEEAAEW

Epitope name EW10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2501)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, optimal epitope, HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- ETINdEAAEW CTL escape mutant elicited a reduced CTL response.

**HXB2 Location** p24 (71–80)

Author Location p24

**Epitope ETINEEAAEW** 

**Immunogen** 

Species (MHC) human (A25)

References Rowland-Jones et al. 1999

• CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied - these women had no delta 32 deletion in CCR5.

- It is in a conserved region, ETINEEAAEW is found in most B, In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive, and the B35 allele seems to be protective.
  - HIV-2 sequence: EIINEEAAEW, no cross-reactivity van Baalen et al. [1996]

**HXB2 Location** p24 (71–80)

Author Location p24 (203–212 SF2)

**Epitope** ETINEEAAEW

Immunogen HIV-1 infection

Species (MHC) human (A25)

Keywords HAART, ART, acute/early infection References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- · The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- · Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A25+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 0/2 group 2, and 1/3 group 3.

**HXB2 Location** p24 (71–80)

**Author Location** p24 (202–211)

**Epitope ETINEEAAEW** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A25)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, CD8 T-cell Elispot granzyme B

**Keywords** Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8+ cells are found, each one constituting 30-40% of the CD8+ cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- 1/3 patients responded to this peptide with GzB producing cells, and the other two responded with IFN-gamma producing cells.

HXB2 Location p24 (71–80)

**Author Location** 

**Epitope DTINEEAAEW** 

Epitope name Gag-DW10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5301)

References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA B\*5301, 2/15 (13%) recognized this epitope.

**HXB2 Location** p24 (71–80)

**Author Location** 

**Epitope** ETINEEAAEW Epitope name Gag-EW10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*5301) References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA B\*5301, 2/15 (13%) recognized this epitope.

**HXB2 Location** p24 (71–80) Author Location p24 (71–80) **Epitope DTINEEAAEW** 

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*5801)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

prediction, immunodominance

References Thakar et al. 2005

- This epitope is highly conserved across clades.
- · PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** p24 (71–80)

**Author Location** p24 (203–212)

Epitope DTINEEAAEW

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (B53)

Keywords HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- · Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B53 women, 0/2 HEPS and 7/9 HIV-1 infected women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in 4 of the 7/9 responsive HIV-1 infected women.

**HXB2 Location** p24 (71–80)

**Author Location** p24 (203–212 subtype A consensus)

**Epitope** DTINEEAAEW

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (B53)

**Keywords** binding affinity, subtype comparisons, epitope processing

References Dorrell et al. 2001

- In clade A infected Gambians, three HLA-B53 epitopes were defined in Gag p24 using ELISPOT, tetramer, and cytotoxicity assays.
- · Two of the new epitopes lacked the predicted P2 anchors, DTI-NEEAAEW and QATQEVKNM, and bound to B53 with high affinity, thus extending the anchor residue motif for B53 and the related B35.
- Two overlapping 20 mer peptides carry this complete epitope, but only one stimulates recognition, which could be due to different peptide processing.
- Keywords subtype comparisons, computational epitope DTINEEAAEW was recognized in only 2/7 HLA-B53 sub-
  - DTINEEAAEW was not A subtype specific and there was cross-recognition although diminished, of the subtype B, C, and D variant, ETINEEAAEW.
  - In one of the two subjects there was cross-recognition of the HIV-2 version of the epitope, EIINEEAADW.

**HXB2 Location** p24 (71–80)

**Author Location** p24

**Epitope** ETINEEAAEW

Subtype A, B, D

Immunogen HIV-1 infection

Species (MHC) human (B53, B58)

Donor MHC A36, A74, B53, B58, Cw4, Cw6; A23, A34, B44, B53, Cw4, Cw6; A23, A24, B35, B58,

Cw4, Cw7;

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, variant cross-

recognition or cross-neutralization References Geels et al. 2005

· Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the

HLA-anchor residues. 42% of the responses were directed to

regions containing new epitopes. This previously described B53 epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype A Gag. The autologous epitope sequence in this person differed from the known epitope by one amino acid, E1D, dTINEEAAEW. It was also recognized in a person carrying a subtype D gag, and in this case the autolo-

gous sequence matched the epitope. This was also predicted to

possibly be a B58 cross-presented epitope in another subtype

B58 motif.

HXB2 Location p24 (71-90)

Author Location p24 (203-222 SF2)

Epitope ETINEEAAEWDRVHPVVHAGP

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A2, B21.

**HXB2 Location** p24 (75–83)

**Author Location** Gag (207–215)

**Epitope** EEAAEWDRV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*40)

**Donor MHC** A\*03, A\*24, B\*35, B\*40

Country United States.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** acute/early infection, variant

recognition or cross-neutralization, superin-

fection

References Yang et al. 2005b

- An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- The response to the epitope EEAAEWDRV was found in the patient before superinfection and diminished afterwards; the initial infecting and superinfecting strain carried EEAAEW-DR1.

HXB2 Location p24 (78-86)

Author Location p24 (78-86)

Epitope AEWDRLHPV

Epitope name AEW

Immunogen HIV-1 infection

Species (MHC) human (A2)

Donor MHC A2, A68.1, B\*07, B\*3503, Cw\*0401,

Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

**Assay type** CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot

- IFNγ

Keywords rate of progression, escape References Oxenius et al. 2004b

D Gag infected person based on peptide reactivity and a known • The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relative efficiency of the variant eptiopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.

This was one of 8 reactive epitopes found not to vary over time.

**HXB2 Location** p24 (78–86)

Author Location Gag (210-218)

Epitope AEWDRVHPV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*40)

**Donor MHC** A\*03, A\*24, B\*35, B\*40

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords acute/early infection, variant cross-

recognition or cross-neutralization, superin-

fection

References Yang et al. 2005b

- An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- The epitope AEWDRVHPV was recognized in the patient before superinfection and diminished afterwards. The initial and superinfecting strains had the variant AEWDR1HPV.

**HXB2 Location** p24 (78–86)

**Author Location** 

Epitope AEWDRVHPV

**Epitope name** Gag-AV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*4002)

**Donor MHC** A\*0201, A\*3201, B\*4002, B\*5301,

Cw\*0202, Cw\*0401

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- · This epitope was newly defined in this study.

- Patient 01RCH59 was Hispanic, not on HAART, and had a viral load of 5100 and CD4 count of 349 – she also recognized TERQANFL, p2p7p1p6(64-71), HLA-B\*4002, and KEKG-GLEGL, Nef(92-100), HLA-B\*4002.
- Among HIV+ individuals who carried HLA B40, 4/5 (80%) recognized this epitope.

HXB2 Location p24 (78–86)
Author Location p24 (78–86)
Epitope AEWDRVHPV
Immunogen HIV-1 infection
Species (MHC) human (B\*4002)
Keywords optimal epitope
References Frahm et al. 2007

HXB2 Location p24 (78–86) Author Location p24 (78–86) Epitope AEWDRLHPV

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*4006)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

- PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.
- >50% conservation across clades.

**HXB2 Location** p24 (81–91) **Author Location** Gag (213–223)

Epitope DRVHPVHAGPI

**Epitope name** Gag 11.4 **Immunogen** vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT, Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining
Keywords subtype comparisons, variant crossrecognition or cross-neutralization, memory
cells

References Amara et al. 2005

 A clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better conserved for CD8 than CD4 T cells.

- All 5 CD8 and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation. The similar reported human epitope in this case is VHPVHAGPIA, restricted by HLA B55.
- 3 of 5 CD8 epitopes and 2 of 8 CD4 epitopes were conserved across multiple HIV-1 clades. DRVHPVHAGPI is identical in HXB2 and the CRF02 consensus. It is relatively conserved across other clades, but usually has an L in the third position: DRIHPVHAGPI.

**HXB2 Location** p24 (81–100)

Author Location p24 (81-100)

Epitope DRLHPVHAGPAAPGQMREPR

Epitope name DRL

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401,

Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

DQ6

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot

- IFNγ

Keywords rate of progression, immunodominance, es-

cape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relative efficiency of the variant epitopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This epitope was not precisely defined, but was one of six epitopes found to be under positive selection for escape mutations and was completely replaced by escape variants between days 66 and 327 (drlhpvhagplapgqmrepr).

**HXB2 Location** p24 (83–92)

**Author Location** p24 (215–223 IIIB)

Epitope VHPVHAGPIA

Immunogen HIV-1 infection Species (MHC) human (B55)

**References** Sipsas *et al.* 1997

- HIV IIIB proteins were used to define the range of CTL epitopes recognized by 3 lab workers accidentally infected with HIV-1 IIIB.
- LHPVHAGPVA, a variant found in HIV-1 PH136, was also recognized.
- LHPVHAGPIA, a variant found in HIV-1 RF, was also recognized.
- LHPVHAGPIT, a variant found in HIV-1 MN, was also recognized.
- LHPAQAGPIA, a variant found in HIV-1 JH3, was recognized at high peptide concentrations.

HXB2 Location p24 (84–91)

Author Location p24

Epitope HPVHAGPI

Subtype D

Immunogen HIV-1 infection Species (MHC) human (B35)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7 **Country** Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

**HXB2 Location** p24 (84–92)

**Author Location** (C consensus)

Epitope HPVHAGPIA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*3910)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- HPVHAGPIA is an optimal epitope.

**HXB2 Location** p24 (84–92)

**Author Location** (C consensus)

Epitope HPVHAGPIA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

HLA class I restricted CD8+ T-cell responses against HIV-1
were analyzed in African patients. Significantly more responses
were shown to be HLA-B restricted. Viral load, CD4 count,
and thus rate of disease progression were also associated with
HLA-B alleles. In addition, the selection pressure imposed on
HIV-1 by HLA-B alleles was shown to be substantially greater
than by other alleles.

• This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (84–92)

**Author Location** (C consensus)

Epitope HPVHAGPIA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- HPVHAGPIA is an optimal epitope.

**HXB2 Location** p24 (84–92)

Author Location p24 (84–92)

Epitope HPVHAGPIA

Epitope name B7-HA9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

Keywords dynamics, supervised treatment interruptions

(STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 1/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection—10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 3/4 individuals had detectable responses to this epitope after STI.

**HXB2 Location** p24 (84–92)

**Author Location** p24 (84–92)

Epitope HPVHAGPIA

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2** Location p24 (84–92)

Author Location p24 (84–92)

**Epitope** HPVHAGPVA

Epitope name B7-HA9 Gag

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** supervised treatment interruptions (STI), es-

cape, early treatment, superinfection

References Altfeld et al. 2002a

- An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.
- The first infecting strain had the variant hpvhagpIa. The CTL response was higher to the second superinfecting variant, HPVHAGPVA.

**HXB2 Location** p24 (84–92)

Author Location (B consensus)

Epitope HPVHAGPVA

Epitope name HA9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A03, B07, Cw7

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
   activation in dying target cells, it was shown that the subset of
   HIV-1-specific CD8+ T cells secreting both IFN-gamma and
   TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1 specific CD8+ T-cell maturation phenotypes and intracellular
   perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** p24 (84–92)

**Author Location** Gag (216–224)

Epitope HPVHAGPIA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

HXB2 Location p24 (87–101)

**Author Location** p24 (219–233 BRU)

Epitope HAGPIAPGQMREPRG

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Claverie et al. 1988

 One of 4 epitopes predicted then shown to stimulate HLA-A2 restricted CTL line.

HXB2 Location p24 (87–101)

Author Location p24 (87–101)

Epitope HAGPIAPGQMREPRG

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 19 patients recognized this epitope.

**HXB2 Location** p24 (87–101)

Author Location Gag (219–233 LAI)

Epitope HAGPIAPGOMREPRG

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Buseyne et al. 1993a

- Vertical transmission of HIV ranges from 13% to 39%.
- Primary assays showed that cytotoxic activity against at least one HIV protein detected in 70% of infected children.
- Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.
- Patient EM28 (CDC P2A) had a CTL response to four epitopes in Gag.

**HXB2 Location** p24 (89–96)

**Author Location** p24 (89–97)

Epitope GPIAPGOM

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*35)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, computational epitope

prediction, immunodominance, optimal epi-

tope

References Thakar et al. 2005

- PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.
- This was a novel epitope.

HXB2 Location p24 (89-96)

Author Location p24

Epitope GPIAPGQM

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (B35)

Donor MHC A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, computational epitope prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

**HXB2 Location** p24 (91–105)

Author Location (C consensus)

Epitope IAPGQMREPRGSDIA

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*13)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression

References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location p24 (91-110)

Author Location p24 (223–242 SF2)

Epitope IAPGQMREPRGSDIAGTTST

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A2, A24, B13, B35.

**HXB2 Location** p24 (93–107)

Author Location Gag (225-239)

Epitope PGQMREPRGSDIAGT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*03, A\*24, B\*35, B\*40

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords acute/early infection, superinfection, charac-

terizing CD8+ T cells

References Yang et al. 2005b

- An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- An early response to this peptide was detected that waned prior to superinfection. The embedded epitope and HLA presenting molecule were not resolved. The initial and superinfecting strains carried a perfect match to the peptide sequence.

HXB2 Location p24 (94–104)

**Author Location** 

Epitope GOMREPRGSDI

Epitope name GI11

Immunogen

Species (MHC) human (B13)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B13 epitope.

HXB2 Location p24 (101-120)

Author Location p24 (233–252 SF2)

Epitope GSDIAGTTSTLQEQIGWMTN

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- · Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A26, A30, B38.

**HXB2 Location** p24 (107–115)

**Author Location** Gag (239–247 SF2)

Epitope TTSTLQEQI

Immunogen vaccine

Vector/Type: Listeria monocytogenes Strain: B clade HXB2 HIV component:

Gag

**Species (MHC)** mouse (H-2K<sup>d</sup>)

**References** Mata *et al.* 1998 LB/c mice were immunized with recombin

- BALB/c mice were immunized with recombinant Listeria monocytogenes (Lm-Gag) expressing HIV-1 HXB2 Gag.
- L. monocytogenes is a gram-positive bacteria that enters the macrophage on phagocytosis and lives in the cytoplasm – secreted L. monocytogenes antigens are processed and presented by both class I and class II pathways.

HXB2 Location p24 (108-117)

**Author Location** 

Epitope TSTLQEQIGW

Subtype B

Immunogen HIV-1 infection **Species (MHC)** human (B\*5701)

Keywords rate of progression, immunodominance

References Migueles & Connors 2001

• HLA B\*5701 was found in a very high frequency in HIV-1 infected non-progressors, 11/13 (85%) versus 19/200 (9.5%) of progressors. Non-progressors tended to have an immune response that was highly focused on four p24 epitopes that were presented by B\*5701, ISPRTLNAW, KAFSPEVIPMF, TSTLQEQIGW, and QASQEVKNW.

**HXB2 Location** p24 (108–117)

**Author Location** 

Epitope TSTLQEQIGW

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*5701)

Keywords rate of progression, immunodominance

References Migueles & Connors 2001

- CTL activity was monitored in 27 individuals, including 10 LTNP with an over-expression of HLA B\*5701 – these individuals have viral loads below the threshold of infection without therapy, and their immune response tends to be focused on peptides that contain B\*5701 epitopes ISPRTLNAW, KAF-SPEVIPMF, TSTLQEQIGW, and QASQEVKNW.
- CTL responses are broader in B\*5701+ individuals with progressive viremia than those that control viremia.
- The HLA-A\*0201 SLYNTVATL epitope response was not as strong in individuals that carried both A2, B57.

**HXB2 Location** p24 (108–117)

**Author Location** 

Epitope TSTLQEQIGN

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Assay type Intracellular cytokine staining, Flow cytomet-

ric T-cell cytokine assay

**Keywords** rate of progression, escape **References** Migueles *et al.* 2003

- cDNA Gag sequences from a set of 17 HLA-B\*5701+ progressors and 10 LTNPs were obtained, and the variation in four p24 B\*5701 epitopes examined. Sequence variants were more common (p < 0.01) in the epitopes in the progressors (median 3, range 1-7) than LTNPs (median 2, range 0-4).</li>
- In general, use of the autologous protein in a target cell did not diminish the overall CD8+ T-cell responses.

**HXB2 Location** p24 (108–117)

**Author Location** (108–117)

**Epitope TSTLQEQIAW** 

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5701, B\*5801)

Assay type Other

Keywords HLA associated polymorphism

**References** Boutwell & Essex 2007

- HIV-1 subtype C proteins were analysed and 94 HLA-associated amino acid polymorphisms were found. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 88% of CTL immunodominant regions had same HLA-associations as found polymorphisms. In 19 studied CTL epitopes, the epitope-restricting allele was same as the polymorphism-restricting allele for 7, and in 12 the alleles differed by specificity.
- TSTLQEQIAW was a previously defined B\*5701/B\*5801 presented epitope that encompassed a B\*57/B\*5801 associated polymorphism, TStLQEQIAW, in the third position.

**HXB2 Location** p24 (108–117)

Author Location (C consensus)

Epitope TSTLQEQIAW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5703)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the T3 and A9 residues of TSTLQE-QIAW are associated with the presence of the HLA presenting molecule in the host.
- TSTLQEQIAW is cross presented by both B\*5801 and B\*5703.

**HXB2 Location** p24 (108–117)

**Author Location** p24 (241–250 LAI)

Epitope TSTVEEQQIW

Subtype B

Immunogen HIV-2 infection

Species (MHC) human (B\*5801)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*5801 epitope.

**HXB2 Location** p24 (108–117)

Author Location p24 (240–249 LAI)

Epitope TSTLQEQIGW

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*5801)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*5801 epitope.

**HXB2 Location** p24 (108–117)

Author Location Gag (240-249)

Epitope TSTVEEQIQW

Epitope name TSTV

Subtype HIV-2

Immunogen HIV-1 infection, HIV-2 infection

Species (MHC) human (B\*5801)

Country Gambia.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Tetramer binding, Flow cytometric T-cell cytokine assay

Keywords characterizing CD8+ T cells

References Gillespie et al. 2005

- CD8 T-cell responses were evaluated and compared in HIV-1 and HIV-2 infected individuals. A significantly greater magnitude and breadth of Gag-specific responses were found in HIV-1 infected individuals, possibly because of undetectable viral load in HIV-2 infected individuals. This finding suggests that responses in HIV-2 infection reflect antigen load in plasma, as is the case in HIV-1 infection. No correlation was found between immune control of HIV-2 and the frequency of perforin-expressing virus-specific CD8 T-cells.
- The HIV-2 epitope TSTVEEQIQW was evaluated in HIV-2 infected individuals, and 4/5 HIV-2-infected B\*5801-positive subjects recognized the TSTV epitope.
- 0/4 HIV-1-infected B\*5801-positive subjects responded to the equivalent TSTLQEQIGW, TSTL epitope.

**HXB2 Location** p24 (108–117)

Author Location Gag (240-249)

Epitope TSTLQEQIGW

Epitope name TSTL

Immunogen HIV-1 infection, HIV-2 infection

Species (MHC) human (B\*5801)

Country Gambia.

Assay type CD8 T-cell Elispot - IFNγ, Tetramer binding, Flow cytometric T-cell cytokine assay

Keywords characterizing CD8+ T cells

References Gillespie et al. 2005

CD8 T-cell responses were evaluated and compared in HIV-1 and HIV-2 infected individuals. A significantly greater magnitude and breadth of Gag-specific responses were found in HIV-1 infected individuals, possibly because of undetectable viral load in HIV-2 infected individuals. This finding suggests that responses in HIV-2 infection reflect antigen load in plasma, as is the case in HIV-1 infection. No correlation was found between immune control of HIV-2 and the frequency of perforin-expressing virus-specific CD8 T-cells.

- The HIV-2 epitope TSTVEEQIQW was evaluated in HIV-2 infected individuals, and 4/5 HIV-2-infected B\*5801-positive subjects recognized the TSTV epitope.
- 0/4 HIV-1-infected B\*5801-positive subjects responded to the equivalent TSTLQEQIGW, TSTL epitope.

**HXB2 Location** p24 (108–117)

Author Location (C consensus)

Epitope TSTLQEQIAW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the T3 residue of TSTLQEQIAW are associated with the presence of the HLA presenting molecule in the host.
- TSTLQEQIAW is cross presented by both B\*5801 and B\*5703.

**HXB2 Location** p24 (108–117)

**Author Location** (C consensus)

Epitope TSTLQEQIAW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5801, B57)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** cross-presentation by different HLA, characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (108–117)

**Author Location** p24 (233–252)

Epitope TSTLQEQIGW

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Bernard et al. 1998

- This study focuses on six rare long-term survivor HIV-infected people who were infected for many years without exhibiting immune dysregulation such immunologically normal HIV-infected (INHI) cases occur at a frequency between 0.1 and 1% in the infected population.
- No direct CTL were found in any of the six INHIs, but above background CTLp activity was found in 3/6 INHIs.
- Epitope sequences were deduced from larger reactive peptides based on HLA binding motifs – XSXXXXXXXW is a B57 binding motif, and CTL activity against TSTLQEQIGW has been found in two other B57 long-term non-progressors.

HXB2 Location p24 (108–117)
Author Location Gag (SF2)
Epitope TSTLQEQIGW
Epitope name TW10
Immunogen HIV-1 infection
Species (MHC) human (B57)

Keywords HAART, ART, acute/early infection

References Goulder et al. 2001a

- Dominant epitope in acute infection in patient PI004, who did not receive any antiviral therapy.
- 1-2 months post seroconversion, subject PI004 displayed a significant decrease in TW10 peptide recognition, followed by an increased CTL response against epitope SL9, SLYNTVATL and other epitopes.
- Three CTL responses, to epitopes TSTLQEQIGW, ISPRTL-NAW, and KAFSPEVIPMF were evident early after infection; CTL responses to SLYNTVATL, QASQEVKNW, EIYKRWII, and FLKEKGGL were detectable at 5 months post-infection and beyond.

HXB2 Location p24 (108–117) Author Location p24 (108–117) Epitope TSTLQEQIGW

Epitope name TST

Immunogen HIV-1 infection Species (MHC) human (B57)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- None of the 8 study subjects recognized this epitope but none were HLA B57+.

HXB2 Location p24 (108–117) Author Location p24 (108–117) Epitope TSTLQEQIGW Immunogen HIV-1 infection

Species (MHC) human (B57)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (108–117)

**Author Location** p24

Epitope TSTLQEQIGW Immunogen HIV-1 infection Species (MHC) human (B57)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

HXB2 Location p24 (108-117)

Author Location p24

Epitope TSTLQEQIGW

Epitope name TST

Immunogen HIV-1 infection Species (MHC) human (B57)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius *et al.* [2000, 2001a] in an IFN? Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there was no correlation between CTL responses with viral rebound rates, plateau viral loads, or clearance rates.

**HXB2 Location** p24 (108–117)

Author Location Gag (147–155)

Epitope TSTLQEQIAW

Epitope name TW10

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B57)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

Keywords epitope processing, escape

References Draenert et al. 2004b

• 174 people who have C clade infections were studied – those who carried B57 have 2 positions in which their HIV Gag consensus is different than the C consensus. One mutation is within this epitope, TW10, at position 3, and is believed to be an anchor residue. The other is in the N-terminal flanking position of the epitope ISPRTLNAW and is thought to impact processing.

HXB2 Location p24 (108–117) Author Location Gag (240–249) Epitope TSTLQEQIAW

Epitope name TW10 Subtype B, C **Immunogen** HIV-1 infection **Species (MHC)** human (B57)

Keywords escape, reversion, viral fitness

References Leslie et al. 2004

• TSTLQEQIAW (the consensus form in the C clade) responses dominate the immune response in HLA-B57 individuals, and this epitope is also recognized in HLA-B5801 individuals. TSnLQEQIAW is shown to be an escape mutant correlated with HLA-B57 and HLA-B5801 alleles. The variant can be transmitted to HLA-B57/B5801 negative individuals, but reverts to wild-type in those. A second escape mutation within the epitope is, however, maintained after transmission; TSNLQE-QIGW is the most common from of the epitope in the B clade, and a G substitution to some other amino acid, often A, was frequently noted in B57+ individuals; transmission of these variants persist in the new host.

HXB2 Location p24 (108–117)
Author Location p24 (108–117)
Epitope TSTLQEQIGW
Immunogen HIV-1 infection
Species (MHC) human (B57)
Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 7 patients recognized this epitope.

**HXB2 Location** p24 (108–117) **Author Location** (B consensus)

Epitope TSTLQEQIGW

Epitope name TW10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

Country United Kingdom.

**Assay type** CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

**Keywords** immunodominance, escape, characterizing CD8+ T cells

References Allen et al. 2004

- This study characterizes an escape mutation in a C-terminal flanking residue of the HLA-A3 gag p17 KK9 epitope that inhibits processing, and is embedded in the overlapping HLA-A3 RK9 epitope.
- The immune response was tracked in subject AC-38. The acute immunodominant response was to the B57 TW10 epitope; this response declined following viral escape (tsNlqeqigw) by day 64. The p17 KK9 and RK9 became immunodominant, but then declined as the escape mutation arose. TW10 is 1 of 3 other strong responses that persisted, along with 1 sub-dominant response.

**HXB2 Location** p24 (108–117)

**Author Location** Gag

Epitope TSTLQEQIGW

Epitope name TW10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Two escape mutations, at positions 3 (T242N) and 9 (G248A), were found in the most polymorphic residues in the epitope. Both were shared between clades B and C. Both were significantly more variable in persons expressing HLA-B57.

**HXB2 Location** p24 (108–117)

Author Location p24 (240-249)

Epitope TSTLQEQIAW

Epitope name TW10

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Ethiopia.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** immunodominance, escape, variant crossrecognition or cross-neutralization

References Currier et al. 2005

- Epitope sequence variation and CD8 T-cell responses were analyzed in C subtype infected HLA-B57-positive individuals from Ethiopia. KF11 was the immunodominant response.
- 9/10 B57 subjects had the T3N TSnLQEQIAW substitution relative to the C consensus, while this form was found in only 2/9 B57- subjects, so it appears to be selected and an immune escape form (p=0.001). Both forms of the TW10 epitope (TSTLQEQIAW and TSnLQEQIAW) were tested in 2 B57-positive subjects; neither responded. The authors suggest this may be due to the dominance of the TW10 response in acute infection, as the response may have been lost by the time of sampling.

**HXB2 Location** p24 (108–117)

**Author Location** Gag

Epitope TSTLQEQIGW

Epitope name TW10

Immunogen HIV-1 infection

Species (MHC) human (B57)

Assay type Other

**Keywords** rate of progression, escape

References Gao et al. 2005

- Three distinct HLA alleles known to alter the rate of AIDS progression were studied. B\*57-mediated protection occurs early in infection and the protective effect of this allele subsides after CD4 cell count drops. In contrast, B\*27 shows no protection against progression to CD4<200, but rather delays progression to an AIDS-defining illness after the CD4 counts have dropped. B\*35-mediated rapid progression to AIDS is probably a function of early decline in CD4 counts.
- TW10 is typically the immunodominant B57 epitope. TW10
  responses are rapid, and escape occurs but presumably with a
  fitness cost, because reversion occurs after the escape variant
  is transmitted to a B57- person. High CD4 counts may be
  maintained in individuals because of immune selection for a
  less fit form of the virus.

HXB2 Location p24 (108–117) Author Location Gag (240–249) Epitope TSTLQEQIAW

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B57)

**Donor MHC** A\*3001, A\*66, B\*4201, B\*5802, Cw\*0602, Cw\*1701; A\*68, A\*66, B\*57, B\*5802, Cw\*0602, Cw\*0602, Cw\*0701

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection

References Pillay et al. 2005

- HIV-specific CTLs in infants were shown to be able to select for viral escape variants early in life, despite a lack of escape with the same CTL specificity in the mother. Infant CTL responses may be compromised by transmission of escape variants that arose in the mother and also those that arose in the father, if the father was the source of the mother's infection.
- TSTLQEQIAW is the C consensus form of the epitope and the autologous form in the mother was TSTLQEQItW, and this was the form transmitted to her infant. The mother does not carry B57, and the B57 escape footprint was inherited paternally. By 33 weeks a new dominant form of the epitope had emerged in the infant, TSnLQEQIAW, and the consensus form was also present in the infant.

**HXB2 Location** p24 (108–117)

**Author Location** p24

Epitope TSTLQEQIGW

Epitope name TW10

Immunogen HIV-1 infection

**Species (MHC)** human (B57, B\*5801)

**Keywords** review, rate of progression, immunodominance, escape, acute/early infection, reversion, viral fitness

References Goulder & Watkins 2004

 This paper is a review of the role of CTL in HIV infection, and it uses the TW10 epitope as an example. HLA B\*57 and B\*5801 both can present this epitope, and are associated with successful containment of HIV infection. The early response to TW10 is immunodominant, and often followed by rapid escape due to the T->N substitution, tsNlqeqigw. Some long term survivors do not carry the escape form, possibly because the CTL response to this epitope is able to suppress viremia. Others do carry the N escape form, and presumably control viremia due to viral attenuation; in support of this the N rapidly back mutates to T in a new host, so there is likely to be a high fitness cost. In contrast, the epitope sometimes contains a G-> A substitution at position 9, and the A can persist in a new host after transmission.

**HXB2 Location** p24 (108–117)

**Author Location** Gag

Epitope TSTLQEQIGW

Epitope name TW10

Immunogen HIV-1 infection Species (MHC) human (B57, B\*5801)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay

**Keywords** responses in children, mother-to-infant transmission, escape

References Feeney et al. 2005

- Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children did not respond to TSTLQE-QIGW but showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.
- TW10 was found to be more frequently recognized by adults than by children. Among B57-positive subjects, TW10 was recognized by 10 out of 12 acutely infected adults, 11/22 chronically infected adults, and only 1/14 infected children. All 14 children carried mutations of this epitope, commonly T3N and G9A (TSnLQEQIaW), but the children were able to recognize the autologous variant. These mutations were rare in adults. One child carried T3N, Q5A, and G9A, and also recognized the autologous variant, TSnLaEQIaW.

**HXB2 Location** p24 (108–117)

**Author Location** p24

Epitope TSTLQEQIGW

Epitope name TW10

Immunogen

**Species (MHC)** (B57, B\*5801)

**Keywords** review, immunodominance, escape, acute/early infection, early-expressed proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

This review discusses the importance of 3 factors that impact
the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA
class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides
listed as immunodominant in acute HIV-1 infection.

**HXB2 Location** p24 (108–117) **Author Location** p24 (235–243)

**Epitope** TSTLQEQIGW

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B57, B58)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

• TSTLQEQIGW cross reacts with both A and B clades.

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** p24 (108–117)

**Author Location** p24 (108–117)

Epitope TSTLQEQIGW Immunogen HIV-1 infection Species (MHC) human (B57, B58)

**Donor MHC** A1, A26, B35, B57, Cw4, Cw\*0601; A1,

A30, B42, B52, Cw7, Cw17; A1, A\*0201,

B44, B57, Cw5, Cw6

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords binding affinity, acute/early infection, early-

expressed proteins, cross-presentation by different HLA

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- This epitope was recognized in three of the acutely infected individuals and was presented by both HLA-B57 and B58.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.

**HXB2 Location** p24 (108–117)

**Author Location** Gag (240–249)

**Epitope** TSTLQEQIGW **Epitope name** gag 240-9

Immunogen HIV-1 infection, HIV-2 infection

Species (MHC) human (B57, B58)

Country Gambia.

Assay type Intracellular cytokine staining

**Keywords** escape, TCR usage, variant cross-recognition or cross-neutralization, characterizing CD8+ T cells

## References Lopes et al. 2003

- CD8+ T cells from HIV-2 infected patients had more polyclonal TCR responses than HIV-1 infected patients, who tended to have oligoclonal responses. This results in limited plasticity of T cell responses to amino acid substitutions within epitopes in HIV-1 infections. HIV-2-specific CD8+ T-cells showed a more diverse TCR usage associated with enhanced CD8 expansion and IFN-gamma production on cross-recognition of variant epitopes.
- This peptide was recognized by a CD8+ T-cell clonotype with Vbeta5.1 usage in one HIV-1 infected patient, and all HIV-1 patients had narrow TCR usage, while HIV-2 patients used multiple TCR Vbeta chains. The HIV-2 variant of this peptide is: tstVEeqiQw. 5/6 HIV-2 infected individuals could recognize both the HIV-1 and HIV-2 peptides, while 0/5 HIV-1 infected patients that could react with the HIV-1 peptide could also react with the HIV-2 peptide.

**HXB2 Location** p24 (108–117)

**Author Location** p24

Epitope TSTLQEQIGW

Subtype B, C

Immunogen HIV-1 infection

**Species (MHC)** human (B57, B58, B63)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This epitope was recognized by 30% of B63-positive subjects and 22% of B57/58-positive subjects.

**HXB2 Location** p24 (108–117)

Author Location p24 (241–250)

Epitope TSTVEEQQIW

Immunogen HIV-2 infection

Species (MHC) human (B58)

References Bertoletti 1998

• HIV-2 epitope defined from an infection in Gambia, Bertoletti,

• All HIV-2 sequences from the database are TSTVEEQIQW in this region, not TSTVEEQQW as in the paper.

**HXB2 Location** p24 (108–117)

Author Location p24

Epitope TSTLQEQIGW

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B58)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied – these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive, and the B35 allele seems to be protective.
- HIV-2 sequence: TSTVEEQIQW, CTL are cross-reactive, Bertoletti et al. [1998]

**HXB2 Location** p24 (108–117) **Author Location** p24 (240–249)

Epitope TSTLQEQIGW Immunogen HIV-2 infection Species (MHC) human (B58)

**Keywords** subtype comparisons, rate of progression, immunodominance

References Bertoletti et al. 1998

- CTL responses in HLA-B\*5801 positive HIV-2 infected individuals have a dominant response to Gag and tolerate extensive substitution, thus HLA-B\*5801+ individuals may have an enhanced potential for cross-protection between HIV-1 and HIV-2.
- This can be an immunodominant epitope in HLA-B57 and B\*5801 infected individuals, and is associated with long-term non-progression Goulder *et al.* [1996b]
- HIV-2 sequence: HIV-2 ROD has the epitope sequence TSTVEEQIQW, and the CTL from a person infected with HIV-2 was cross-reactive with HIV-1 epitopes.
- The epitope is TSTLQEQIGW in HIV-1 B clade, and TSTVEE-QIQW in HIV-2 ROD.
- HLA B\*5801 and B35 may preferentially select HIV-1 and HIV-2 cross-reactive epitopes.

**HXB2 Location** p24 (108–117) **Author Location** p24 (240–249 SF2)

Epitope TSTLQEQIGW
Immunogen HIV-1 infection
Species (MHC) human (B58)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B58+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 1/1 group 2, and 0/0 group 3.

HXB2 Location p24 (108–117) Author Location p24 (108–117) Epitope TSTLQEQIGW Epitope name TW10 Immunogen HIV-1 infection Species (MHC) human (B58)

**Keywords** acute/early infection **References** Goulder *et al.* 2001c

- Responses to this dominant A3-restricted Gag epitope are present during the time of decreasing viral load in acute infection.
- Mutations in this epitope were observed in autologous clones of subjects who were B58-positive with a higher frequency than those who were B58-negative (P = 0.02)
- These mutations are being sexually transmitted in adult infections.

**HXB2 Location** p24 (108–117)

**Author Location** 

**Epitope** TSTLQRQIGW **Immunogen** HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls (ML1250).

**HXB2 Location** p24 (108–117)

Author Location Gag (B57)

Epitope TSNLQEQIGW

Epitope name TW10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Characteristic changes in B57 epitopes in B57+ people were mapped: TSNLQEQIGW often has one or both of the substitutions: tsNlqeqygw, tstlqeqy[A/G]w.

**HXB2 Location** p24 (108–118)

**Author Location** p24 (240–249 LAI)

Epitope TSTLQEQIGWF

Subtype B

Immunogen HIV-1 infection
Species (MHC) human (B\*57, B\*5801)
Keywords rate of progression
References Goulder *et al.* 1996b

- Response to this epitope was found in 4 slow progressing HLA-B\*57 individuals, in 2 it was dominant or very strong.
- For one donor (from Zimbabwe) this was defined as the optimal peptide.
- This epitope can be presented in the context of the closely related HLA molecules B\*5801 and B\*57.

HXB2 Location p24 (108–118) Author Location p24 (240–249 LAI) Epitope TSTLQEQIGWF Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*5701) Keywords optimal epitope References Frahm *et al.* 2007

• C. Brander notes this is a B\*5701 epitope.

**HXB2 Location** p24 (108–118)

**Author Location** 

**Epitope** TSTLQEQIGWF **Epitope name** Gag-TF11

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57) References Sabbaj *et al.* 2003

 Among HIV+ individuals who carried HLA B57, 2/5 (40%) recognized this epitope.

HXB2 Location p24 (109–117) Author Location Gag (241–249 LAI) Epitope STLQEQIGW

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5701, B\*5801)

**Keywords** rate of progression **References** Klein *et al.* 1998

- B57 has been associated with long-term non-progression in the Amsterdam cohort.
- The most pronounced CTL responses in HLA B\*5701 LTS were to RT and Gag.

**HXB2 Location** p24 (109–117)

**Author Location** 

**Epitope** STLQEQIGW **Epitope name** Gag-SW9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57) References Sabbaj *et al.* 2003

- Among HIV+ individuals who carried HLA B57, 1/5 (20%) recognized this epitope.
- Among HIV+ individuals who carried HLA B58, 1/4 (25%) recognized this epitope.

**HXB2 Location** p24 (109–117) **Author Location** p24

Epitope STLQEQIGW Subtype B, D

Immunogen HIV-1 infection Species (MHC) human (B58)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7 **Country** Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The epitope sequence in this person had a single G8A change, STLQEQIaW.

HXB2 Location p24 (110–118) Author Location Gag (242–) Epitope TLQEQIGWM

Epitope name Gag242

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: p24 Gag Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was an intermediate A2 binder that did not induce CTL and CD8+ T-cell IFN gamma responses in mice.
   Responses were detected in 2/17 HIV+ HLA-A2 subjects.

**HXB2 Location** p24 (118–126)

Author Location Gag

Epitope MTSNPPIPV

**Epitope name** Gag 271

Subtype M

Immunogen vaccine, in vitro stimulation or selection

Vector/Type: DNA, peptide Adjuvant: In-

complete Freund's Adjuvant (IFA)

**Species (MHC)** human, mouse (A\*0201)

Assay type Cytokine production, T-cell Elispot

**Keywords** subtype comparisons, computational epitope prediction, escape, TCR usage, variant cross-recognition or cross-neutralization

References McKinney et al. 2004

- This study examined variant recognition of epitopes presented by A\*0201 and A\*1101. Numerous amino acid substitutions can be introduced into epitopes without disrupting their recognition by CTLs, although epitopes with multiple substitutions were less recognized. An algorithm was constructed for prediction of epitopes capable of inducing responses to a great number of variant epitopes.
- MTSNPPIPV form is most common in subtype C while MTnNPPIPV form is mostly found in subtype B.
- A total of 14 variant forms of Gag 271 were identified. Immunization with MTSNPPIPV form induced CTLs recognizing 11 of the variant forms while MTnNPPIPV form induced CTLs recognizing only 3 of the epitope variants.

HXB2 Location p24 (118–126) Author Location p24 (118–126) Epitope MTNNPPIPV

Epitope name MV9

Immunogen HIV-1 infection Species (MHC) human (A2)

**Donor MHC** A\*02, A\*23, B\*07, B\*51, Cw\*15

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection, reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs.
   Escape variants were commonly detected in maternal plasma.
   Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- The mother was A02- and carried a variant form of the epitope, MThNPPIPV, which she passed to her A02+ child. This form persisted in her child for 12 months.

HXB2 Location p24 (118–126)
Author Location Gag (282–290)
Epitope MTNNPPIPV
Immunogen HIV-1 infection
Species (MHC) human (A2 supertype)
Keywords supertype, rate of progression
References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.

- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind three of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

HXB2 Location p24 (121–135)
Author Location p24 (253–267)
Epitope NPPIPVGEIYKRWII
Immunogen HIV-1 infection
Species (MHC) human (B8)

References Gotch et al. 1990

• High frequency of memory and effector Gag-specific CTL.

HXB2 Location p24 (121–135)

Author Location p24 (255–274 SF2)

Epitope NPPIPVGEIYKRWII

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Keywords** review, immunodominance, escape **References** Goulder *et al.* 1997a; Phillips *et al.* 1991

- Longitudinal study of CTL escape mutants in people with the appropriate HLA types – little variation was observed in the immunodominant B27 epitope, relative to B8 epitopes, which varied over time.
- Goulder et al. [1997a] is a review of immune escape that points out that there may be a protective effect associated with B27, and that HLA-B8 individuals tend to progress more rapidly than HLA B27 patients.

HXB2 Location p24 (121–135)

Author Location p24 (121–135)

Epitope NPPIPVGEIYKRWII

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location p24 (121–135)

Author Location p24 (121–135 HXB2)

Epitope NPPIPVGEIYKRWII

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment in

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

 Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.

- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 19% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p24 (121–140) Author Location p24 (253-272)

Epitope NPPIPVGEIYKRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

• HIV-specific CTL lines developed by ex vivo stimulation with peptide.

**HXB2 Location** p24 (121–140)

**Author Location** p24 (253–272 SF2)

Epitope NPPIPVGEIYKRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- protein.
- · Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- Two of these 12 had CTL response to this peptide.
- The responding subjects were HLA-A2, A3, B8, B62, and HLA-A1, B8, B18.

HXB2 Location p24 (121-140)

Author Location p24 (253–272 SF2)

Epitope NPPIPGEIKRWIILGNIK

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

• CTL expanded *ex vivo* were later infused into HIV-1 infected

**HXB2 Location** p24 (121–140)

**Author Location** p24 (255–274 SF2)

Epitope NPPIPVGEIYKRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human

References van Baalen et al. 1993

• Gag CTL epitope precursor frequencies were estimated and peptide mapping was performed.

HXB2 Location p24 (121-142)

**Author Location** p24 (253–274 BH10)

Epitope NPPIPVGEIYKRWIILGLNKIV

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Johnson et al. 1991

Gag CTL response studied in three individuals.

HXB2 Location p24 (121-152)

**Author Location** Gag

Epitope NPPIPVGEIYKRWIILGLNKIVRMYSPTSILD

Immunogen HIV-1 infection, vaccine

Vector/Type: lipopeptide HIV component:

Gag

Species (MHC) human (A\*0201)

References Seth et al. 2000

- Immunization of 2/4 HIV seropositive HLA selected individuals with a 32 amino acid Gag lipopeptide that contains CTL epitopes restricted by HLA A33, B8, B27, B35, and Bw62 gave a transient increase in peptide-specific bulk CTL response, but they did not decrease plasma viral load.
- Placebo and HLA mis-matched controls showed no change in
- The responders carried HLA Bw62 and B35 the two HLAmatched that did not respond carried B35 and B8.

**HXB2 Location** p24 (121–152)

Author Location Gag (183–214 LAI)

Epitope NPPIPVGEIYKRWIILGLNKIVRMYSPTSILD

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Of 25 patients, most had CTL specific for more than 1 HIV-1 Anti-HIV lipopeptide vaccine consisting of six long amino acid peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I
  - A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees - 9/10 reacted to this peptide.
  - 9/12 tested mounted a CTL response to at least one of the six peptides; each of the six peptides elicited a CTL response in at least one individual - this peptide was particularly immunogenic, eliciting a CTL response in four vaccinees.
  - All of the 12 tested had an IgG response to this peptide.

**HXB2 Location** p24 (122–130)

**Author Location** (C consensus)

**Epitope** PPIPVGDIY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*35)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 Defined as minimal peptide by titration curve, PPIPVGEIY individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the D7 residue of PPIPVGDIY are associated with the presence of the HLA presenting molecule in the host.

HXB2 Location p24 (122-130) **Author Location** (C consensus)

Epitope PPVPVGDIY

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*35)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- PPVPVGDIY is an optimal epitope.

HXB2 Location p24 (122-130)

Author Location p24 (260-268 LAI)

**Epitope** PPIPVGDIY

Subtype B

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human (B\*3501)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*3501 epitope.

HXB2 Location p24 (122-130)

Author Location p24 (245–253 HIV-2)

Epitope NPVPVGNIY

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

References Rowland-Jones et al. 1995

HXB2 Location p24 (122-130)

Author Location p24 (245–253 HIV-2)

Epitope NPVPVGNIY

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*3501 epitope.

HXB2 Location p24 (122-130)

Author Location p24 (260–268 LAI)

Epitope PPIPVGDIY

Subtype B

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human (B35)

References Rowland-Jones et al. 1995

and HIV-2 form NPVPVGNIY are also recognized.

HXB2 Location p24 (122-130)

Author Location p24 (260-268 LAI)

Epitope PPIPVGDIY

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (B35)

References Lalvani et al. 1997

- · A peptide-based protocol was optimized for restimulation of CTLp using optimized peptide and IL-7 concentrations - importantly this protocol does not stimulate a primary response, only secondary - peptide-specific CTLp counts could be obtained via staining with peptide-Class I tetramers.
- This peptide was one of the B35 presented test peptides used in control experiments showing that the assay gave no activity using lymphocytes from 21 healthy B35 seronegative donors.

HXB2 Location p24 (122-130)

Author Location p24 (260-268 LAI)

**Epitope** PPIPVGDIY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords review

References McMichael & Walker 1994

• Review of HIV CTL epitopes.

HXB2 Location p24 (122-130)

**Author Location** p24 (subtype B)

Epitope PPIPVGEIY

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B35)

Keywords subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi - these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- · Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found - B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among B and D clade viruses.
- The Clade A version of the epitope, PPIPVGDIY, was preferentially recognized by CTL.

HXB2 Location p24 (122-130)

**Author Location** 

**Epitope PPIPVGEIY** 

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords acute/early infection References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL re- Patient SC15 (HLA A1/68, B8/35, Bw4/6, Cw4/0704) was sponses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects -3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- · No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

HXB2 Location p24 (122-130)

**Author Location** p24

Epitope PPIPVGDIY

Immunogen

Species (MHC) human (B35)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied - these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive, and the B35 allele seems to be protective.
- HIV-2 version of this epitope is not conserved: NPVPVGNIY, but the CTLs are cross-reactive – one of five B35 CTL epitopes that are cross-reactive, see also Rowland-Jones et al. [1995]

**HXB2 Location** p24 (122–130)

**Author Location** p24 (260–268)

Epitope PPIPVGDIY

Epitope name PPI

Immunogen HIV-1 infection Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- One of two HLA B35+ among the eight study subjects recognized this epitope.

given acute and sustained therapy and recognized epitopes PPIPVGDIY and VPLRPMTY during 331 days of HAART treatment.

HXB2 Location p24 (122–130)

Author Location p24 (122-130)

Epitope PPIPVGDIY

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Ferrari et al. 2000

One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2** Location p24 (122–130)

**Author Location** p24 (254–262 SF2)

Epitope PPIPVGDIY

**Immunogen** HIV-1 infection

Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B35+ individuals that had a CTL response to this epitope broken down by group: 0/2 group 1, 0/2 group 2, and 1/1 group 3.

HXB2 Location p24 (122-130)

Author Location p24 (260–268)

Epitope PPIPVGDIY

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B35)

Keywords HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.

- Among HLA-B35 women, 1/3 HEPS and 3/4 HIV-1 infected women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in the 1/3 HEPS case and in the all 3/4 responsive HIV-1 infected women.
- Subject ML 857 shifted from a A\*6802 DTVLEDINL and B35 (H/N)PDIVIYQY response prior to seroconversion to a B35 PPIPVGDIY and B35 VPLRPMTY response postseroconversion.

**HXB2 Location** p24 (122–130)

**Author Location** 

**Epitope** PPIPVGDIY **Epitope name** Gag-PY9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35) References Sabbaj *et al.* 2003

- Among HIV+ individuals who carried HLA B35, 2/21 (10%) recognized this epitope.
- Among HIV+ individuals who carried HLA B\*5301, 0/11 (0%) recognized this epitope.

**HXB2 Location** p24 (122–130)

Author Location p24

Epitope PPIPVGEIY Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

*Vector/Type:* DNA prime with modified vaccinia Ankara (MVA) boost *Strain:* A clade *HIV component:* p17 Gag, p24 Gag

HIV component. p17 Gag, p24

Species (MHC) human (B35)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

**HXB2 Location** p24 (122–130) **Author Location** p24 (260–268)

Epitope PPIPVGDIY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot granzyme B

Keywords characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- None of three patients responded to this peptide with GzB producing cells and one of the patients responded with IFNgamma producing cells.

**HXB2 Location** p24 (122–130)

**Author Location** Gag

**Epitope** PPIPVGEIY **Immunogen** HIV-1 infection

Species (MHC) human (B35)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** HIV exposed persistently seronegative (HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- No one, 0/3 HLA B35+ infection-resistant men, and 0/5 preseroconversion men who went on to become infected, reacted to this epitope.

HXB2 Location p24 (122-130)

Author Location p24 (122-130)

Epitope PPIPVGDIY

Immunogen HIV-1 infection

Species (MHC) human (B35)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses.
   HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 4/9 patients recognized this epitope.

HXB2 Location p24 (122–130)

**Author Location** (C consensus)

Epitope PPVPVGDIY

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B35) Country South Africa.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$  **Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (122–130)

Author Location p24

**Epitope PPIPVGEIY** 

Subtype B, D

Immunogen HIV-1 infection

Species (MHC) human (B35, B53)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons

**References** Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence in this person matched the known epitope. In another D subtype infected individual, it was predicted to be a B53 epitope based on HLA typing of the individual and motifs within the reactive peptide.

**HXB2 Location** p24 (122–130)

**Author Location** Gag (254–262)

**Epitope PPIPVGEIY** 

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (B7 supertype)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

HXB2 Location p24 (122-130)

Author Location p24

Epitope PPIPVGDIH

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls, ML887.

HXB2 Location p24 (124–138)

Author Location p24 (256–270 LAI)

Epitope IPVGEIYKRWIILGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Buseyne et al. 1993b

 Clustering of Gag p24 CTL epitopes recognized in 29 HIVinfected people.

**HXB2 Location** p24 (124–138)

Author Location Gag (256–270 LAI)

Epitope IPVGEIYKRWIILGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Buseyne et al. 1993a

- Vertical transmission of HIV ranges from 13% to 39%.
- Primary assays showed that cytotoxic activity against at least one HIV protein was detected in 70% of infected children.
- Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.
- Two children, EM16 (CDC P2A+D2) and EM18 (CDC P2A), had a CTL response to this epitope, and it was shown to be presented by B8 in EM18.

**HXB2** Location p24 (126–140)

**Author Location** p24 (126–140 HXB2)

Epitope VGEIYKRWIIGLNK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%).
   p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 19% of the study subjects, and it was one of the 25 most frequently recognized peptides.

HXB2 Location p24 (127–135)

Author Location p24 (259–267 SF2)
Epitope GDIYKRWII
Immunogen HIV-1 infection
Species (MHC) human (B\*0801)
References McAdam et al. 1998

GDIYKRWII specific CTL clone also recognized GEIYKR-WII

**HXB2 Location** p24 (127–135)

**Author Location** p24 (261–269)

Epitope GEIYKRWII
Immunogen HIV-1 infection

Species (MHC) human (B8)

References Sutton et al. 1993

 Predicted epitope based on B8-binding motifs, from larger peptide NPPIPVGEIYKRWII.

HXB2 Location p24 (127-135)

**Author Location** p24 (259–267)

Epitope GEIYKRWII

Immunogen in vitro stimulation or selection

Species (MHC) human (B8)

Keywords dendritic cells

References Zarling *et al.* 1999

- This study compares the ability of macrophages and dendritic cells to stimulate primary responses in CD8+ lymphocytes isolated from HLA-appropriate HIV-uninfected donors using peptide-pulsed APC – the dendritic cells performed better as APC for the stimulation of primary responses.
- Strong CTL responses were elicited by the epitopes DRFYK-TLRA and GEIYKRWII when presented by either immature or mature dendritic cells – macrophages were not able to prime a CTL response against DRFYKTLRA.
- A weak response to KLTPLCVSL was stimulated using macrophages as the APC.
- No detectable response was observed for the following previously-defined HIV epitopes: KIRLRPGGK, ILKEPVHGV, IRLRPGGK, GPKVKQWPL.

**HXB2 Location** p24 (127–135)

Author Location p24 (259–267 LAI)

Epitope GEIYKRWII

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Klenerman et al. 1994

Naturally occurring variant GDIYKRWII may act as antagonist.

**HXB2 Location** p24 (127–135)

**Author Location** p24 (259–267)

Epitope GEIYKRWII

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Keywords** immunodominance **References** Betts *et al.* 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the HLA A2+ was HLA A\*0201, A31, B8, B51 and responded to this epitope as well as seven others.

**HXB2 Location** p24 (127–135)

**Author Location** p24 (259–267)

Epitope GEIYKRWII

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords dynamics, escape

References Nowak et al. 1995

 Longitudinal study of CTL response and study of immune escape – GDIYKRWII could also stimulate CTL, reactivity fluctuated.

**HXB2 Location** p24 (127–135)

Author Location p24 (259–267)

Epitope GEIYKRWII

Immunogen HIV-1 infection

Species (MHC) human (B8)

**References** McAdam *et al.* 1995

 Equivalent sequence GDIYKRWII also recognized by CTL from some donors. **HXB2 Location** p24 (127–135)

**Author Location** p24 (259–267)

Epitope GEIYKRWII

Epitope name GEI

Immunogen HIV-1 infection Species (MHC) human (B8)

**Keywords** HAART, ART, supervised treatment interruptions (STI), immunodominance, escape, acute/early infection

## References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- Six of the 7/8 study subjects that were HLA B8 recognized this epitope.
- Patient SC2 (HLA A1, B7/8, Cw0701/0702, DR4/53, DQ7)
  had CTL responsiveness against epitopes FLKEKGGL, GPKVKQWPL, and GEIYKRWII peptides FLKEKGGL tetramer staining steadily declined and at day 1340 the FLKEKGGL
  stained cells were no longer detected and the escape mutant
  FLKENGGI was found in 8/10 clones.
- Patient SC9 (HLA A1/2, B8/13, Cw0/0701, DR2/11, DQ6/7) had a CTL response against epitopes FLKEKGGL, ILKEPVHGV, SQRRQDILDLWIYHTQGYFPDWQNY, and GEIYKRWII and all responses declined during therapy initiated at day 390 but were restored when therapy become intermittent.
- Patient SC19(HLA A11/12, B8/44, Cw06/0701, DR3/7, DR52/53, DQ 2/8) had a CTL response to epitopes FLKEKGGL, GEIYKRWII, ACQGVGGPGHK, AVDLSH-FLK, and FNCGGEFFY that declined during therapy initiated at day 197.
- Patient SC10(HLA A1/3, B8/35, DR1/8, DQ 4/5) had sustained therapy started during acute infection and maintained an immunodominant response to FLKEKGGL and a response to GEIYKRWII through day 1088.
- Patient SC12(HLA A1, B8/39, Cw0701/0702, DR2/3, DR51/52, DQ2/6) had sustained therapy started during acute infection and maintained an immunodominant response to FLKEKGGL throughout and minor responses to GEIYKR-WII, DCKTILKAL, GGKKKYKLK – GEIYKRWII and GGKKKYKLK responses were stimulated by a brief period off therapy.
- Patient SC11(HLA A1, B8, Cw0201, DR3/11, DR52, DQ2/7) started therapy early, remained on therapy for 40 days, then reinitiated HAART at day 640 had a CTL response to FLKEKGGL, GPKVKQWPL, and GEIYKRWII throughout and received a benefit from the early limited course therapy.

**HXB2 Location** p24 (127–135)

Author Location p24 (259–267 SF2)

Epitope GEIYKRWII Immunogen HIV-1 infection Species (MHC) human (B8)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B8+ individuals that had a CTL response to this epitope broken down by group: 2/3 group 1, 2/3 group 2, and 2/2 group 3.

HXB2 Location p24 (127-135)

Author Location p24

Epitope GEIYKRWII

Epitope name GEI

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius et al. [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there
  was no correlation between CTL responses with viral rebound
  rates, plateau viral loads, or clearance rates.

HXB2 Location p24 (127–135)

Author Location p24

**Epitope** GEIYKRWII **Subtype** A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

Species (MHC) human (B8)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

• The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].

 Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location p24 (127–135) Author Location Gag (259–267) Epitope GEIYKRWII Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (B8)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes, variant cross-recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the most frequently recognized of the 31 epitopes that the vaccinated volunteers responded to.

HXB2 Location p24 (127–135) Author Location p24 (259–267) Epitope GEIYKRWII Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8) Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- None of three patients responded to this peptide with GzB producing cells and one of the patients responded with IFNgamma producing cells.

**HXB2 Location** p24 (127–135)

Author Location p24

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Epitope GEIYKRWII Subtype B, D

Immunogen HIV-1 infection Species (MHC) human (B8)

**Donor MHC** A1, A1, B8, B55, Cw3, Cw7 **Country** Democratic Republic of the Congo. **Assay type** CD8 T-cell Elispot - IFNγ **Keywords** subtype comparisons **References** Geels *et al.* 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence matched the peptide.

**HXB2 Location** p24 (127–135)

Author Location Gag (259–267 BRU)

**Epitope** GEIYKRWII **Subtype** B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (B8)
Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 1/9 CRF02\_AG-infected Ivorians, and 2/9 B-infected French subjects.

**HXB2 Location** p24 (127–136)

**Author Location** 

Epitope GEIYKRWIIL

**Epitope name** Gag-GL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

**Donor MHC** A\*0101 A\*0301 B\*0801 B\*5802 Cw\*0602 Cw\*0701

Assay type Chromium-release assay

**Keywords** HAART, ART **References** Sabbaj *et al.* 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- Among HIV+ individuals who carried HLA B08, 3/6 (50%) recognized this epitope.

**HXB2 Location** p24 (128–135) **Author Location** p24 (260–267 LAI)

Epitope EIYKRWII Subtype B

Immunogen

Species (MHC) human (B\*0801)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*0801 epitope.

**HXB2 Location** p24 (128–135)

**Author Location** (C consensus)

Epitope DIYKRWII

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (128–135)

Author Location (C consensus)

Epitope DIYKRWII

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- DIYKRWII is an optimal epitope.

**HXB2 Location** p24 (128–135)

Author Location p24 (260–267 LAI)

Epitope EIYKRWII

Subtype B

Immunogen

Species (MHC) human (B8)

References Goulder et al. 1997g

• Defined in a study of the B8 binding motif.

**HXB2 Location** p24 (128–135)

**Author Location** p24 (SF2)

Epitope EIYKRWII

**Immunogen** HIV-1 infection

Species (MHC) human (B8)

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in an HIV+ Caucasian living in Boston – this epitope did not fall within the three most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

HXB2 Location p24 (128–135)

Author Location p24 (C consensus)

Epitope DIYKRWII

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in an HIV+ South African – this epitope did not fall within the five most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

**HXB2 Location** p24 (128–135)

Author Location p24 (SF2)

Epitope EIYKRWII

Epitope name EI8

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Goulder et al. 2001a

- This peptide elicited a weak CTL response during acute HIV-1 infection in patient PI004.
- Three CTL responses to epitopes, TSTLQEQIGW, ISPRTL-NAW, and KAFSPEVIPMF, were evident early after infection; CTL responses to SLYNTVATL, QASQEVKNW, EIYKRWII, and FLKEKGGL were detectable at 5 months post-infection and beyond.

HXB2 Location p24 (128–135)

Author Location p24

Epitope EIYKRWII
Immunogen HIV-1 infection
Species (MHC) human (B8)
Keywords rate of progression
References Kostense *et al.* 2001

- HLA tetramers to six epitopes were used to study HLA-A2, B8 and B57 CTL in 54 patients HIV-specific tetramer positive cells were inversely correlated with viral load in patients with high CD4, but in patients with CD4 T-cells below 400 high tetramer frequencies were found despite high viral load.
- Most patients have high levels of HIV-specific T-cell expansions, but many of these cells aren't functional.
- In 15 of the patients, the proportion of IFN gamma producing tetramer cells correlated with AIDS-free survival.
- 4/13 patients that reacted with EIYKRWII displayed epitope mutations in a minority of sequences, which did not correlate with disease progression or viral load – these mutations were: Patient 156 (KIYKRWMI), Patient 36 (EIYKRRII), Patient 656 (KIYKRWII, EIYERWMI), and Patient 159 (EIYKRWVI).
- Stimulation with HLA-B8 p24 and Nef epitopes significantly increased Nef-specific T-cell numbers in 2 patients (748 and 1113)
- There were more functional IFN-gamma producing Nefspecific T-cells within the T-cell population than there were active p24 Gag-specific T-cells.

HXB2 Location p24 (128–135) Author Location p24 (259–267) Epitope DIYKRWII Immunogen HIV-1 infection Species (MHC) human (B8) References Appay et al. 2000

- Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virus-specific CD8+ T cells produced IFN- $\gamma$  and MIP-1 $\beta$  with a distinct subset that failed to produce TNF- $\alpha$

HXB2 Location p24 (128–135)
Author Location p24 (128–135)
Epitope EIYKRWII
Immunogen HIV-1 infection
Species (MHC) human (B8)
References Day et al. 2001

• B8-restricted CTL accounted for about 1/3 of the total CTL response in one individual.

HXB2 Location p24 (128–135)

Author Location Gag

Epitope EIYKRWII

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Goulder et al. 2000b

- Tetramer assays were compared with three functional assays in 42 people with chronic HIV infection: ELISPOT, intracellular cytokine staining, and precursor frequency (limiting dilution assay [LDA])
- HIV-specific tetramer staining CTLs appeared to be active, and inert CTL were not found to play a significant role in chronic pediatric or adult HIV infection.

HXB2 Location p24 (128–135)
Author Location p24
Epitope DIYKRWII
Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A2, A11, B8, B60, Bw6 **Keywords** HAART, ART

References Appay *et al.* 2002
• Four HIV patients with prolonged clinically successful antiviral therapy but with ongoing evidence of replication and Nef mRNA transcription, showed specific T-cell responses by

Elispot and Tetramer staining, maintained for 2-4 years after initiation of HAART.

Nef epitope recognition was detected in all 4 subjects, gp120,
 Pol and Gag-specific in 1 or 2 subjects.

 The HIV-specific CD8+ T-cells had an intermediate maturation phenotype characterized by low levels of perforin and high levels of CD27 expression.

HXB2 Location p24 (128–135) Author Location Gag (260–267 IIIB)

 ${\bf Epitope} \ {\tt EIYKRWII}$ 

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B8)

Assay type Chromium-release assay

References Kurane et al. 2003

Three CD8+ CTL cell clones were derived from 2 HIV-1 positive asymptomatic patients, and their epitope specificities and HLA presenting proteins were defined.

HXB2 Location p24 (128–135) Author Location Gag (B con) Epitope EIYKRWII Epitope name E18 Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8)

**Country** United States. **Assay type** CD8 T-cell Elispot - IFNγ

Keywords variant cross-recognition or cross-

neutralization

References Draenert et al. 2004c

• CD8+ T-cell responses that persisted in individuals with advanced disease were studied; most of the optimal epitopes defined were recognized with intermediate to high avidity. On average 13 (range, 2-39) epitopic regions were targeted in an average of 6 proteins (range, 1-8). HAART resulted in decrease in antigen and reduction in gamma IFN EliSpot responses, suggesting active responses to autologous virus. The lack of mutations within most viral epitopes suggest that persistent CTL through late disease do not exert strong immune selection

pressure, yet the Elispot assays show robust responses, suggesting to the authors that gamma IFN-based screening methods may not reveal functional CD8+ T-cell impairment in patients with AIDS.

 3 subjects recognized this epitope with intermediate functional avidity. Autologous sequence revealed one substitution, Diyrkwil, in 1 of the 3; this version of the epitope also had intermediate functional avidity with the donor's cells.

HXB2 Location p24 (128-135)

Author Location Gag

Epitope EIYKRWII
Immunogen HIV-1 infection
Species (MHC) human (B8)
Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** HIV exposed persistently seronegative (HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- 3/9 HLA B8+ infection-resistant men, compared to 1/3 preseroconversion men who went on to become infected, reacted to this epitope.

**HXB2 Location** p24 (128–135)

Author Location (B consensus)

Epitope EIYKRWII

Epitope name EI8

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8)

**Donor MHC** A02, A03, B08, B62, Cw7, Cw10; A11, A29,

B08, B44, Cw4, Cw7; A25, A32, B08, B14, Cw7, Cw8; A01, A03, B08, B14, Cw7, Cw8

Country United States.

Assay type Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 4/9 individuals recognized this epitope, presented by HLA-B8.

**HXB2 Location** p24 (128–135) **Author Location** p24 Epitope DIYKRWII

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B8)

Country United Kingdom.

Assay type Tetramer binding, T-cell Elispot, Intracellular

cytokine staining

**Keywords** rate of progression, acute/early infection, characterizing CD8+ T cells, immune dysfunction

References Papagno et al. 2004

Acute HIV-1 infection induces massive activation of HIV-specific and non-HIV-specific CD8+ T-cells resulting in differentiation of these cells. High differentiation of CD8+ T-cells is correlated with disease progression. Differentiation is a natural process but it can be driven by elevated immune activation, such as in HIV infection.

HXB2 Location p24 (128–135)

Author Location p24 (128-135)

Epitope EIYKRWII

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A1, A3, B8, B35

Country United States.

**Assay type** Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, escape, variant cross-recognition or cross-neutralization

References Casazza et al. 2005

- Evidence of continued viral evolution during HAART therapy despite low viral load was found in 1/5 patients studied.
- Diykrwii sequence was found in 12/17 clones after the initiation of therapy, while eVykrwii sequence was found in 5/17, and the peptide used to initially detect the response was not found, EIYKRWII. The less frequent clone was most often recognized. No dramatic shift towards escape was observed after the initiation of therapy.

HXB2 Location p24 (128-135)

**Author Location** Gag

Epitope EIYRKWII

Immunogen HIV-1 infection

Species (MHC) human (B8)

Country Netherlands.

Assay type Tetramer binding, Flow cytometric T-cell cy-

tokine assay

Keywords binding affinity, rate of progression, escape,

characterizing CD8+ T cells

References Jansen et al. 2005

Number and responsiveness of CD8 T-cells directed to different Gag peptides presented by HLA-A2, -B8 and B57 were compared. It was shown that T-cells specific for an HLA-B57 peptide responded to a higher extent and more readily to antigenic stimulation than those specific for HLA-A2 and -B8 peptides did. Moreover, it was shown that the higher functionality of B57-restricted T-cells was not correlated to higher number of epitope escape mutations in A2- and B8-restricted T-cells.

- Tetramer decay experiments indicate that the HLA-B57 peptide has a higher half-life than the A2 and B8 peptides. The authors point out that CD8+ T cells with high binding affinity may require less help.
- In 1/2 B8+ individuals that were sequenced, 2 epitope variants were present: EfYRKWII and rIYRKWII, but the form EIYRKWII was found most often.

HXB2 Location p24 (128-135)

Author Location Gag (260–267 B consensus)

Epitope EIYKRWII

Epitope name EI8

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8)

**Donor MHC** A\*01, A\*11, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** escape, variant cross-recognition or crossneutralization, optimal epitope

References Koibuchi et al. 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- The EI8 variant dIYKRWII was the only form of the epitope detected over a 5 year time period in this person. Elispot reactions were comparable between the autologous form and the B clade consensus form, EIYKRWII.

HXB2 Location p24 (128-136)

**Author Location** Gag (260–268 SUMA)

**Epitope** EIYKRWIIL **Epitope name** Gag EIL9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*2402)

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, acute/early infection, characterizing CD8+ T cells

References Jones et al. 2004

• Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.

- The patient SUMA maintained low viral loads and stable CD4
   T cell counts through seven years of follow up. In contrast
   to more rapid progressors, WEAU and BORI, SUMA a broad
   response to 24 epitopes, with little immunodominance. Two
   peptides were somewhat more intensely recognized in acute
   infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

**HXB2 Location** p24 (128–136)

Author Location Gag (260-268)

Epitope EIYKRWIIL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (A2)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

**Keywords** vaccine-induced epitopes, variant cross-

recognition or cross-neutralization **References** Gahéry-Ségard *et al.* 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the most frequently recognized of the 31 epitopes that the vaccinated volunteers responded to.

**HXB2 Location** p24 (128–136)

Author Location p24

Epitope EIYKRWIIL

Subtype B, D

Immunogen HIV-1 infection

Species (MHC) human (A24)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence in this person matched the known epitope.

HXB2 Location p24 (129–136)
Author Location p24 (263–270 SF2)
Epitope IYKRWIIL
Immunogen HIV-1 infection
Species (MHC) human (A\*2402)
References Ikeda-Moore et al. 1997

- Defined using reverse immunogenetics 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) 53 of the 59 peptides bound A\*2402.
- This peptide induced CTL in 1/4 HIV-1 + people tested.
- IYKRWIIL bound to A\*2402 with medium strength, the epitope can be processed in a vaccinia construct and presented two specific CTL clones were obtained.

HXB2 Location p24 (129–138)
Author Location p24 (263–272 SF2)
Epitope IYKRWIILGL
Immunogen HIV-1 infection
Species (MHC) human (A\*2402)
References Ikeda-Moore et al. 1997

- Defined using reverse immunogenetics 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) 53 of the 59 peptides bound A\*2402.
- This peptide induced CTL in 1/4 HIV-1 + people tested.
- IYKRWIILGL bound to A\*2402 with medium strength, the epitope can be processed in a vaccinia construct and presented two specific CTL clones were obtained.

HXB2 Location p24 (129–138) Author Location Gag (261–270) Epitope IYKRWIILGL Subtype B

Immunogen vaccine

Vector/Type: lipop

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (A24)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

• After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

**HXB2 Location** p24 (129–138)

Author Location p24

 ${\bf Epitope} \ {\tt IYKRWIILGL}$ 

Subtype B, D

Immunogen HIV-1 infection Species (MHC) human (A24)

Donor MHC A23, A24, B35, B58, Cw4, Cw7

**Country** Democratic Republic of the Congo.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$  **Keywords** subtype comparisons **References** Geels *et al.* 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence in this person matched the known epitope.

Author Location p24 (129–138)
Author Location p24 (263–272)
Epitope IYKRWIILGL
Immunogen HIV-1 infection
Species (MHC) human (B27)
Keywords immunodominance

**References** Betts *et al.* 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals was B27 and responded to IYKR-WIILGL.

**HXB2 Location** p24 (129–148) **Author Location** Gag (261–280)

Epitope IYKLWIILGLNKIVRMYSPT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27, B62) **Donor MHC** A3, A31, B27, B38; A24, B27, B62

Assay type Chromium-release assay

Very and conital and mysesslimmunit

Keywords genital and mucosal immunity

References Musey et al. 2003

- CTL clones from blood, semen, cervix and rectum samples from 12 individuals were compared. CTL clones derived from blood and mucosal samples had similar high lysis efficiency, were primarily perforin dependent, and TCRbeta VDJ region sequencing revealed cases of expansion of the same clone in different compartments.
- CD8+ T cell clones directed at this epitope were derived from blood and rectum of one individual, and blood and semen of another. Both individuals are HLA-B27 positive, and within the peptide there is a B27 epitope that was recognied in the blood and rectum of the first patient, and in the blood of the second. A HLA-B62 epitope is also recognized in this peptide in the second individual, and the CD8+ T cells clones from both the blood and semen recognized this epitope.

**HXB2 Location** p24 (130–148) Author Location p24 (265–280 BRU) Epitope YKRWIILGLNKIVRMYSPT

Immunogen HIV-1 infection Species (MHC) human (B27) References Dadaglio et al. 1991

• Used as a positive control for HLA specificity.

**HXB2 Location** p24 (131–139) Author Location Gag (265-273) Epitope KRWIILGLN Immunogen HIV-1 infection **Species (MHC)** chimpanzee (Patr-B\*03)

References Balla-Jhagjhoorsingh et al. 1999b

- · Certain HLA-alleles have been associated with long-term survival – among them are HLA-B\*27 and HLA-B\*57.
- Of more than 150 chimpanzees that have been reported to be infected with HIV-1, only one has developed AIDS.
- CTL responses were studied in two HIV-1 infected chimpanzees that have strong CTL responses, and they were found to respond to highly conserved epitopes that are recognized in humans in the context of HLA-B\*27 and HLA-B\*57.
- The human HLA protein which presents this Patr-B\*03 epitope is HLA B\*2705 but the amino acid sequences in the binding pockets of HLA-B\*2705 and Patr-B\*03 are distinctive.

**HXB2 Location** p24 (131–140) Author Location p24 (263–272) Epitope KRWIILGLNK Immunogen HIV-1 infection Species (MHC) human (B\*27) Keywords HAART, ART References Huang et al. 2000

- · The single cell ELISPOT assay was optimized and highly specific, and found to work well even after the primary cells had been frozen and thawed.
- Increases in gamma interferon producing cells were observed in response to anti-retroviral therapy using single cell IFNgamma-production ELISPOT.
- In 3/3 HLA A\*02, B\*27 individuals, the dominant response in gag measured by both gamma IFN production and T cell lysis was to the B27 epitope, KRWIILLGLNK, not the A2 SLYNTVATL epitope.

HXB2 Location p24 (131-140) Author Location p24 (263-272 SF2) Epitope KRWIILGLNK Immunogen HIV-1 infection Species (MHC) human (B\*27) References McAdam et al. 1998

• Epitope invariant across clades A, B, C, and D.

**HXB2 Location** p24 (131–140) Author Location p24 (260–269 HIV-2) Epitope RRWIQLGLQK Immunogen

Species (MHC) human (B\*2703) Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is a B\*2703 epitope.

**HXB2 Location** p24 (131–140)

Author Location p24

Epitope KRWIILGGLNK Immunogen HIV-1 infection Species (MHC) human (B\*2705)

Keywords dynamics, acute/early infection

References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers – high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- Tetramers with peptide variants KRWIILGGLNK and KRWI-IMGGLNK were used – CTL from most B27 donors recognize both variants, although one of the three subjects recognized only KRWIILGGLNK.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects – 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- · No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

**HXB2 Location** p24 (131–140) Author Location p24 (263–272 LAI)

**Epitope** KRWIILGLNK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*2705)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*2705 epitope.

**HXB2 Location** p24 (131–140)

**Author Location** p24 (263–272)

**Epitope** KRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human (B\*2705)

Keywords escape

References Kelleher et al. 2001b

- A mutation in 4/5 B\*2705 patients had substitution to lysine (K) at HIV-1 gag residue 264 (R264K), in three the change occurred late in infection - in one patient a substitution of glycine at HIV-1 gag residue 264 (R264G) was detected - these substitutions reduce binding to B27.
- The R264K mutations were associated with a L268M mutation that may be compensatory, and R264G occurred in conjunction with E260D.

- Positions 260, 264, and 268 all lie along one aspect of helix seven of the capsid protein, a region that is important for capsid self-association and assembly.
- R264G and R264K escape mutation outgrowth occurred in conjunction with high viral loads.

HXB2 Location p24 (131–140) Author Location p24 (263–272) Epitope KRWIIMGLNK Immunogen HIV-1 infection Species (MHC) human (B\*2705) References Appay *et al.* 2000

- Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virusspecific CD8+ T cells produced IFN- $\gamma$  and MIP-1 $\beta$  with a distinct subset that failed to produce TNF- $\alpha$

**HXB2 Location** p24 (131–140)

Author Location p24

**Epitope** KRWIILGLNK **Subtype** A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade

HIV component: p17 Gag, p24 Gag

Species (MHC) human (B\*2705)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the A-clade epidemic in Nairobi, Kenya. A DNA and MVA prime-boost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location p24 (131–140) Author Location Gag (263–272) Epitope KRWIILGLNK Epitope name KK10 Subtype B

Immunogen HIV-1 infection, vaccine

Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag, gp120, gp41, Protease

Species (MHC) human (B\*2705)

**Donor MHC** A2, B27, B44, Cw2, Cw5

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Tetramer binding,

Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay, Degranulation, CD107a and b cell sur-

face mobilization

**Keywords** vaccine-specific epitope characteristics, vaccine-induced epitopes, immunodominance, escape, TCR usage, characterizing

CD8+ T cells

References Betts et al. 2005

- A vaccinated HIV-negative individual exhibited features predicted to be necessary for vaccine-induced protection: killing ability, cytokine production, multifunctionality, proliferative capacity, polyclonality, memory phenotype, and a CD8 T-cell response directed against a highly immunodominant epitope correlated with long-term nonprogression. In spite of this, the subject became infected with HIV through homosexual contact. The subject progressed more rapidly than expected for an HLA-B27-positive individual.
- After infection, CD4+ and CD8+ T cells acquired functional characteristics typical of chronic HIV infection. The infecting virus escaped the vaccine-induced T-cell response with an R264G substitution, KgWIILGLNK, which diminishes binding to B27, between the second and third year of infection.

**HXB2 Location** p24 (131–140)

Author Location p24 (263–272 LAI)

Epitope KRWIILGLNK

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*2705, B27)

**Keywords** review, rate of progression, immunodominance, escape

References Goulder et al. 1997c; Goulder et al. 1997a

- HLA-B\*2705 is associated with slow HIV disease progression.
- 11/11 HLA-B\*2705 donors make a response to this epitope, usually an immunodominant response.
- This is a highly conserved epitope.
- The HLA-B\*2705 binding motif includes R at position 2, and L in the C-term position.
- Goulder et al. [1997a] is a review on CTL immune escape that discusses this epitope in the context of the difficulty in detection of immune escape KRWIILGLNK and an R2K change, KKWIILGLNK, show little difference in titration curves, yet the K2 variants fail to bind to targets for more than 1 hour, while the R2 form can sensitize lysis by CTL for over 24 hours minigene transfection experiments confirmed the importance of this for the CTL response.

HXB2 Location p24 (131–140) Author Location p24 (260–269 HIV-2) Epitope RRWIQLGLQK Immunogen

Species (MHC) human (B27)

References Brander & Walker 1996

• HIV-2, HLA-B\*2703, S. Rowland-Jones, pers. comm.

HXB2 Location p24 (131-140)

Author Location p24 (263-272 LAI)

Epitope KRWIILGLNK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords dendritic cells

References Fan et al. 1997

 The capacity of dendritic cells to process and present antigen and stimulate anti-HIV-1 CTL memory responses was studied.

**HXB2 Location** p24 (131–140)

**Author Location** Gag (263–272)

Epitope KRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords epitope processing, dendritic cells

References Zheng et al. 1999

- Protein delivery (gp160 LAV, p66 LAV, and p24 NY5) to human dendritic cells (DC) with liposomes provides enhanced memory CTL response relative to delivery of protein alone.
- Chloroquine administration enhanced epitope presentation, and brefeldin A and peptide aldehyde inhibitors inhibited antigen presentation, suggesting epitopes were processed by classical proteasome pathway.
- The CTL response to p24 was measured in individuals with a response to B27-KRWIILGLNK.

**HXB2 Location** p24 (131–140)

Author Location p24 (263–272 LAI)

Epitope KRWIILGLNK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords dynamics, TCR usage

References Wilson et al. 1998a

- HIV+ individuals were followed longitudinally using MHC tetramers in combination with 14 anti-BV chain MAbs, and clonal expansion of HIV-specific T cells was followed *in vivo*.
- Seven HIV+ people were studied, and all showed expansions of particular TCR BV clones, often several, relative to uninfected controls.
- Three patients were followed in detail, TCR VB expansions persisted for 2 to 3 years, with occasional transient increases.

**HXB2 Location** p24 (131–140)

**Author Location** p24

Epitope KRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords review

References Rowland-Jones et al. 1997

• Described in this review as the first identified HIV CTL epitope.

HXB2 Location p24 (131-140)

**Author Location** p24 (263–272 LAI)

Epitope KRWIILGLNK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Buseyne et al. 1993b

 Clustering of Gag p24 CTL epitopes recognized in 29 HIVinfected people.

**HXB2 Location** p24 (131–140)

Author Location p24 (263-272 LAI)

Epitope KRWIILGLNK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords review

References McMichael & Walker 1994

• Review of HIV CTL epitopes.

**HXB2 Location** p24 (131–140)

**Author Location** p24 (263–272)

Epitope KRWIIMGLNK

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Klenerman et al. 1994

Naturally occurring variant KRWIILGLNK may act as antagonist.

**HXB2 Location** p24 (131–140)

Author Location p24 (263–272)

Epitope KRWIIMGLNK

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Klenerman et al. 1995

Naturally occurring variant KRWIILGLNK may act as antagonist

**HXB2 Location** p24 (131–140)

Author Location p24 (265–274)

Epitope KRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords dynamics, TCR usage

References Moss et al. 1995

- In one individual, TCR usage changed over time indicating that new populations of CTL can be recruited.
- TCR usage showed a CTL clonal response to this epitope that persisted over 5 years.
- CTL clones specific for HIV epitopes may represent between 0.2 and 1% of the total CD8+ population of T cells.

**HXB2 Location** p24 (131–140)

Author Location p24 (265–276)

Epitope KRWIILGLNK

Immunogen

Species (MHC) human (B27)

References Carreno et al. 1992

• Included in HLA-B27 binding peptide competition study.

HXB2 Location p24 (131-140)

Author Location p24 (265–274 SF2)

Epitope KRWIILGLNK Immunogen HIV-1 infection Species (MHC) human (B27)

**Keywords** dynamics, review, immunodominance, escape **References** Goulder *et al.* 1997a; Phillips *et al.* 1991

- Longitudinal study of CTL escape mutants little variation was observed in the immunodominant B27 epitope, relative to B8 epitope.
- Goulder *et al.* [1997a] is a review of immune escape that points out that there may be a protective effect associated with B27, and that HLA-B8 individuals tend to progress more rapidly than HLA B27 patients.

HXB2 Location p24 (131–140) Author Location p24 (263–272) Epitope KRWIILGLNK Immunogen HIV-1 infection Species (MHC) human (B27)

**Keywords** review, escape

References Goulder et al. 1997a; Nietfeld et al. 1995

- Single point mutations were introduced and viral viability and CTL recognition tested – an Arg to Lys change at anchor position P2 abrogates binding to B27, but doesn't change viral viability *in vitro*.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study.

HXB2 Location p24 (131–140)
Author Location p24 (263–272)
Epitope KRWIIMGNK
Immunogen HIV-1 infection
Species (MHC) human (B27)

Keywords escape

References Nowak et al. 1995

 Longitudinal study of CTL response and immune escape – the form KRWIILGNK was also found, and both forms stimulate CTL.

HXB2 Location p24 (131–140)
Author Location p24 (263–272)
Epitope KRWIILGNK
Immunogen HIV-1 infection
Species (MHC) human (B27)
Keywords subtype comparisons
References Durali et al. 1998

antigens expressed in vaccinia.

- Cross-clade CTL response was studied by determining the CTL activity in seven patients from Bangui, (6 A subtype, and 1 AG recombinant infections) and one A subtype infection from a person living in France originally from Togo, to different
- Pol reactivity: 8/8 had CTL to A subtype, and 7/8 to B subtype, and HIV-2 Pol was not tested.
- Gag reactivity: 7/8 reacted with A or B subtype gag, 3/8 with HIV-2 Gag.
- Nef reactivity: 7/8 reacted with A subtype, and 5/8 with B subtype, none with HIV-2 Nef.
- Env reactivity: 3/8 reacted with A subtype, 1/8 with B subtype, none with HIV-2 Env.
- One of the patients was shown to react to this epitope: KRWI-ILGNK.

HXB2 Location p24 (131–140) Author Location p24 (263–272) Epitope KRWIIMGLNK Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords review, immunodominance, escape

References Goulder et al. 1997f; Goulder et al. 1997a

- Six HLA-B27 donors studied make a strong response to this epitope.
- In 4/6 cases, this was the immunodominant or only CTL response.
- Two of the cases had an epitope switch to the form KKWI-IMGLNK during a period of rapid decline to AIDS, following their asymptomatic period.
- The arginine to lysine switch is in an anchor residue, and results in immune escape due to severely diminished binding to the B27 molecule.
- Goulder *et al.* [1997a] is a review of immune escape that summarizes this study in the context of CTL escape to fixation.

**HXB2 Location** p24 (131–140)

Author Location p24

Epitope KRWIILGLNK

Immunogen

Species (MHC) human (B27)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied – these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive, and the B35 allele seems to be protective.
- HIV-2 sequence: RRWIQLGLQK this epitope was not HIV-1 and HIV-2 cross-reactive.

**HXB2 Location** p24 (131–140)

**Author Location** Gag (263–)

Epitope KRWIILGLNK

Immunogen computer prediction

Species (MHC) (B27)

**Keywords** subtype comparisons

References Schafer et al. 1998

- This study uses EpiMatrix for T cell epitope prediction to identify possible HLA-B27 and A-2 CTL epitopes in HIV.
- Based on EpiMatrix predictions, 28 peptides were synthesized and tested using T2 binding assays for potential HLA A2 or B27 binding, and 12 of these were shown to bind to the predicted HLA molecule.
- Two of these 12 peptides had been previously identified as CTL epitopes: HLA-B27 KRWILGLNK and HLA-A2 ILKEPVHGV.
- This peptide sequence is not conserved between clades, but is found in most B clade isolates.

**HXB2 Location** p24 (131–140)

**Author Location** p24 (263–282)

Epitope KRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Bernard et al. 1998

- This study focuses on six rare long-term survivor HIV-infected people who were infected for many years without exhibiting immune dysregulation - such immunologically normal HIVinfected (INHI) cases occur at a frequency between 0.1 and 1% in the infected population.
- No direct CTL were found in any of the six INHIs, but above background CTLp activity was founded in 3/6 INHIs.
- Epitope sequences were deduced from larger reactive peptides based on HLA binding motifs - XRXXXXXXX is a B\*2705 binding motif.

HXB2 Location p24 (131-140)

Author Location p24 (265–274 SF2)

Epitope KRWIILGLNK Immunogen HIV-1 infection Species (MHC) human (B27)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- · Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- · The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B27+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 0/0 group 2, and 1/1 group 3.

HXB2 Location p24 (131-140)

Author Location p24 (263–272)

Epitope KRWIILGLNK

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (B27)

Keywords HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- · Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Subject ML 1760 had an A2 response to ILK(D/E)PVHGV prior to seroconversion, and gained responses to epitopes A2 SL(F/Y)NTVATL and B27 KRWII(L/M)GLNK postseroconversion.

**HXB2 Location** p24 (131–140)

**Author Location** p24 (131–140)

Epitope KRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Day et al. 2001

**HXB2 Location** p24 (131–140)

**Author Location** p24 (260–299)

Epitope RRWIQLGLQK

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Day et al. 2001

**HXB2 Location** p24 (131–140)

**Author Location** p24 (131–140)

**Epitope** KRWIILGLNK

Epitope name KK10

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords responses in children, mother-to-infant transmission, immunodominance, escape,

acute/early infection

References Goulder et al. 2001b

- 85% of B27+ adults have CTL that recognize this epitope, but only 2/6 children did.
- Responses to this dominant B27-restricted Gag epitope are present during the time of decreasing viral load in acute infec-
- Three children who shared B27 with their mothers did not respond to this epitope and inherited escape mutations from their mothers.
- · A transmitted R132T anchor residue mutation abrogated binding to B27.
- In the three children infected with the non-binding KK10 variants, the dominant CTL specificity was still HLA-B27restricted, but it was directed against an epitope in p17, IRL-RPGGKK, only rarely recognized in adults when KRWIIL-GLNK is the dominant response.
- Mutations in this epitope were observed in autologous clones of subjects who were B27-positive with a higher frequency than those who were B27-negative (P = 0.0005)
- These mutations are being sexually transmitted in adult infec-

HXB2 Location p24 (131-140)

**Author Location** 

**Epitope** KRWIILGLNK

Epitope name Gag-KK10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Sabbaj et al. 2003

Among HIV+ individuals who carried HLA B27, 2/3 (66%) recognized this epitope.

**HXB2 Location** p24 (131–140)

Author Location p24 (263–272 LAI)

**Epitope** KRWIIMGLNK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B27)

Keywords HAART, ART, epitope processing, immun-

odominance

References Kelleher et al. 2001a

- Ritonavir (RTV) inhibits chymotryptic activity in the 20S proteasome *in vitro*, as does Saquinavir (SQV) to a lesser extent; Indinavir (IDV) does not. Thus there is concern protease inhibitors may adversely effect CTL epitope processing, but this paper indicates that processing is not inhibited at therapeutically relevant concentrations of RTV when the proteasome is functioning in an intracellular context.
- RTV did not reduce antigen presentation and concentration of the two immunodominant Gag CTL epitopes (KRWIIMGLNK (B27) and SLYNTVATL (A2)).
- RTV did not inhibit the processing and assembly of HLA-B35 or -A2, which are assembled with a rapid and moderate time course, respectively, or of HLA-A3, -B27 and -B39.

**HXB2 Location** p24 (131–140)

Author Location p24

 ${\bf Epitope} \ {\sf KRWIILGLNK}$ 

Epitope name B27-KK10(p24)

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B27)

Donor MHC A24, B7, B27; A30, A32, B18, B27

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals. Patient C displayed the greatest response to B27-KK10(p24), and in decreasing order also responded to A24-RW8(Nef), B7-IL9(gp41), A24-RL9(gp41), A24-YL8(gp41), and B7-TM9(Nef). Patient D also displayed the greatest response to B27-KK10(p24), and also responded to A30-RY11(p17), A32-PW10(RT), A30-KY11(RT), A32-RW10(gp120), and B18-YY9(Nef).

**HXB2 Location** p24 (131–140)

Author Location Gag (263–272)
Epitope KRWIILGLNK

**Subtype** CRF01\_AE **Immunogen** HIV-1 infection

Species (MHC) human (B27)

Donor MHC B27

**Keywords** subtype comparisons **References** Currier *et al.* 2002a

- Cross-reactive responses were found in PBMC isolated from individuals infected with either B or CRF01\_AE clade viruses, as determined by Elispot assays of target cells expressing recombinant vaccinia viruses expressing HIV-1 gag, env, nef and pol from many clades.
- Subject AIHP-6 (Thai, CDF01-AE infected) recognized this
  epitope. This subject showed cross-subtype CTL responses to
  gag constructs derived from subtypes A, B, C, D, F, G, and
  H, and this epitope was perfectly preserved in all of these but
  subtype A which had the sequence KRWMILGLNK.
- This subject didn't respond to a Gag CRF01 sequence which had a R->K mutation in position 2.

HXB2 Location p24 (131-140)

**Author Location** Gag

Epitope KRWIILGLNK

Epitope name KK10

Immunogen HIV-1 infection

Species (MHC) human (B27)

Donor MHC A26, B27

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords responses in children, rate of progression, im-

munodominance, escape **References** Feeney *et al.* 2004

- Viral load in a perinatally infected child remained low until emergence of an escape variant (kTwiilglnk) in the immunodominant CTL epitope KRWIILGLNK when the child was 7.4 years old. The emergence of this escape mutation was followed by an increase in viremia and an increase in the number of targeted CTL epitopes, measured again when the child was 9.2 years old. The timing suggests that the loss of recognition of this epitope may have resulted in the subsequent loss of
- The mutation krwillMglnk has been suggested to be compensatory and required for the emergence of the previously described escape mutation kKwillMglnk (Kelleher 2001). The L136M mutation does appear in this child, but not in conjunction with the R132T escape mutation.

**HXB2 Location** p24 (131–140)

**Author Location** 

immune control.

Epitope KRWIIMGLNK

Epitope name KK10

Immunogen HIV-1 infection Species (MHC) human (B27)

**Keywords** review, responses in children, vaccine-specific

epitope characteristics, rate of progression,

escape

References Goulder & Watkins 2004

- This paper is a review of the role of CTL in HIV infection, and it uses KK10 as an example of an epitope that has late escape mutations associated with loss of immune control of the virus and decline to AIDS.
- A study where a vaccine response to KK10 was stimulated in a individual who subsequently got infected and had rapid escape from the KK10 response and an unexpectedly high steady state viral load for a B27+ person is recounted as a cautionary note regarding the delicate balance of effects that might contribute to CTL mediated immune control.

**HXB2 Location** p24 (131–140)

**Author Location** Gag

Epitope KRWIILGLNK

Epitope name KK10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B27)

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 6,krwiiMglnk, was found in the most polymorphic residue in the epitope. This was shared between clades B and C. One escape mutation, at position 2, kTwiilglnk, was found not to correspond to the most polymorphic residue in the epitope.

**HXB2 Location** p24 (131–140)

**Author Location** Gag (263–272 BRU)

**Epitope** KRWIILGLNK Subtype B, CRF02 AG Immunogen HIV-1 infection

Species (MHC) human (B27)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ **Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 0/9 CRF02\_AG-infected Ivorians, and 3/9 B-infected French subjects.

HXB2 Location p24 (131-140)

**Author Location** p24

**Epitope** KRWIILGLNK

Epitope name KK10

Immunogen

Species (MHC) (B27)

Keywords review, immunodominance, escape, acute/early infection. early-expressed proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

• This review discusses the importance of 3 factors that impact the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides listed as immunodominant in acute HIV-1 infection.

HXB2 Location p24 (131–140)

Author Location Gag (263-272)

Epitope KRWIILGLNK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

Country Barbados.

Assay type CD8 T-cell Elispot - IFNγ

Keywords responses in children, mother-to-infant trans-

mission, escape, acute/early infection

References Pillay et al. 2005

- HIV-specific CTLs in infants were shown to be able to select for viral escape variants early in life, despite a lack of escape with the same CTL specificity in the mother. Infant CTL responses may be compromised by transmission of escape variants that arose in the mother and also those that arose in the father, if the father was the source of the mother's infection.
- A B27 escape footprint was passed from a B27+ father to his partner, who then passed the variant to their child. KRWIIL-GLNK is the B consensus form of this epitope, the paternal form was KqWIIiGLNK, the maternal form, KqWvImGLNK, and this was the form passed on to the child.

**HXB2 Location** p24 (131–140)

Author Location Gag (263–272)

Epitope KRWIILGLNK

Immunogen HIV-1 infection

Species (MHC) human (B27)

Country Australia.

Assay type CD8 T-cell Elispot - IFNγ, HLA binding

**Keywords** binding affinity, rate of progression, immun-

odominance, escape

References Ammaranond et al. 2005

• B27-positive subjects have an immunodominant response to Gag epitope KRWIILGLNK, with R264 KrWIILGLNK being a crucial anchor residue. Among a group of 20 long-term non-progressive B27-positive subjects, 14 carried wild-type sequence, 5 carried known escape mutants (K264 or G264), and 1 carried a novel Q264 mutant. This mutant was also shown to be a likely escape mutation. These escape mutations all lower the affinity for B27 binding; the Q264 variant has 30-fold lower binding affinity.

**HXB2 Location** p24 (131–140)

Author Location Gag (263–272 LAI)

Epitope KRWILLGLNK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Buseyne et al. 1993a

- Vertical transmission of HIV ranges from 13% to 39%.
- · Primary assays showed that cytotoxic activity against at least one HIV protein was detected in 70% of infected children.
- · Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.
- Patient EM28 (CDC P2A) had a CTL response to four epitopes in Gag.

**HXB2 Location** p24 (131–142)

**Author Location** p24 (265–276)

Epitope KRWIILGLNKIV

Immunogen peptide-HLA interaction

Species (MHC) human (B27)

References Jardetzky et al. 1991

B27.

**HXB2 Location** p24 (131–142)

Author Location p24 (263-274 LAI)

Epitope KRWIILGLNKIV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords dendritic cells

References Fan et al. 1997

• The capacity of dendritic cells to process and present antigen and stimulate anti-HIV-1 CTL memory responses was studied.

**HXB2 Location** p24 (131–142)

Author Location p24 (131–142)

Epitope KRWIILGLNKIV

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location p24 (131-145)

Author Location p24 (263–277 LAI)

Epitope KRWIILGLNKIVMRY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A33)

**References** Buseyne et al. 1993b

• Clustering of Gag p24 CTL epitopes recognized in 29 HIVinfected people.

**HXB2 Location** p24 (131–145)

**Author Location** p24 (266–277)

Epitope KRWIILGLNKIVRMY

Immunogen vaccine

Vector/Type: vaccinia HIV component: Gag

Species (MHC) human (B27)

References Nixon et al. 1988

- Gag CTL epitope mapped with rec gag-vaccinia and synthetic
- This was the first HIV-1 epitope to be mapped.

**HXB2 Location** p24 (131–145)

**Author Location** p24 (266–277 LAI)

Epitope KRWIILGLNKIVMRY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Meyerhans et al. 1991

Longitudinal study showing persistence of epitope despite CTL

**HXB2 Location** p24 (131–145)

Author Location p24 (265–279)

Epitope KRWIILGLNKIVRMY

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Nixon et al. 1990; Rowland-Jones et al. 1999

- Epitope examined in the context of peptide binding to HLA- HIV-1 and HIV-2 cross-reactive CTL clone, highly conserved
  - · Reviewed in Rowland-Jones99, notes that it did not appear cross-reactive with HIV-2 in Rowland-Jones98, HIV-2 form: RRWIQLGLQK.

**HXB2** Location p24 (131–145)

Author Location p24 (SF2)

Epitope KRWILGLNKIVRMY

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in a HIV+ African American living in Boston with unknown HLA - this epitope did not fall within the three most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

**HXB2 Location** p24 (131–145)

Author Location p24 (131–145 HXB2)

Epitope KRWIILGLNKIVRMY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

• Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.

- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- Responses to this peptide were detected in 16% of the study subjects, and it was one of the 25 most frequently recognized peptides.

HXB2 Location p24 (131–146)
Author Location p24 (265–279)
Epitope KRWIILGLNKIVRMYC
Immunogen HIV-1 infection
Species (MHC) human (B27)
References Bouillot et al. 1989

 HLA-B27 restricted epitope also binds to HLA-A2 and HLA-B37 in solid phase assay.

HXB2 Location p24 (131–150)

Author Location p24 (265–284 SF2)

Epitope KRWIILGLNKIVRMYSPTSI

Immunogen HIV-1 infection

Species (MHC) human (B62?)

References van Baalen et al. 1993

• Gag CTL epitope precursor frequencies estimated.

HXB2 Location p24 (131–150)
Author Location p24 (263–282 SF2)
Epitope KRWIILGLNKIVRMYSPTSI
Immunogen HIV-1 infection
Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 A-2 had CTL response to this peptide.
- The responding subject was HLA-A3, A32, B51, B62.

HXB2 Location p24 (131–152)

Author Location p24 (263–284 BH10)

Epitope KRWIILGLNKIVRMYSPTSILD

Immunogen HIV-1 infection

Species (MHC) human (B62)

References Johnson et al. 1991

• Gag CTL response studied in three individuals.

HXB2 Location p24 (132–140) Author Location Gag (261–280) Epitope RWIILGLNK Subtype B Immunogen HIV-1 infection Species (MHC) human (B27) **Donor MHC** A24, A33, B14, B27; A2, A32, B27, B62 **Assay type** Chromium-release assay **Keywords** genital and mucosal immunity **References** Musey *et al.* 2003

- CTL clones from blood, semen, cervix and rectum samples from 12 individuals were compared. CTL clones derived from blood and mucosal samples had similar high lysis efficiency, primarily perforin dependent, and TCRbeta VDJ region sequencing revealed cases of expansion of the same clone in different compartments.
- CD8+ T cell clones that recognize this epitope were derived from both blood and cervix from a woman, and the blood and semen from a man.

HXB2 Location p24 (132–145)

Author Location Gag
Epitope KWILGLNKIVRMY
Immunogen HIV-1 infection

Species (MHC) human (B27)
Keywords TCR usage
References Weekes et al. 1999b

- Peptide 728: Almost all CD8+ T cells are CD28+ at birth, and the proportion of CD28-CD8+ cells increases with age – this study examines the contribution of CD8+CD28- cells to CTL memory pools for CTL clones specific for two persistent human viruses, CMV and HIV – clones were found to be similarly distributed the CD28 depleted cell population.
- HIV CTL responses to 3 Env and 2 Gag peptides were studied.
- The clonal composition of the TCR  $V\beta$  responses were studied and was found to be highly focused, with one TCR beta-chain sequence tending to dominate the peptide-specific response clones to this epitope were  $V\beta$ 22.1.

HXB2 Location p24 (132–145)

Author Location Gag
Epitope KWILGLNKIVRMY
Immunogen HIV-1 infection
Species (MHC) human
References Weekes et al. 1999a

 Peptide 728: Memory CTL specific for HIV-1 may contribute to oligoclonal expansions within the CD57+ CD28- CD8+ CTLp populations.

Author Location p24 (134–143)
Author Location Gag (266–275)
Epitope IILGLNKIVR
Subtype B
Immunogen vaccine
Vector/Type: lip.

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (A3, A11)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

• After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

**HXB2 Location** p24 (134–143) **Author Location** p24 (subtype B) **Epitope IILGLNKIVR** Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A33)

**Keywords** subtype comparisons References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi - these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found - B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among A, B and D clade viruses.

**HXB2 Location** p24 (135–142) **Author Location** p24 (135–142) Epitope ILGLNKIV Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

- PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.
- This epitope is highly conserved across clades.

HXB2 Location p24 (135–142) Author Location Gag (267-274) **Epitope ILGLNKIV** Subtype B Immunogen vaccine Vector/Type: lipopeptide Strain: B clade

LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (A2)

Assay type proliferation, CD8 T-cell Elispot - IFNγ, Chromium-release assay

Keywords vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

• After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the most frequently recognized of the 31 epitopes that the vaccinated volunteers responded to.

**HXB2 Location** p24 (135–142) Author Location Gag (267–274 HXB2)

> Epitope ILGLNKIV Subtype B, CRF01\_AE

Immunogen HIV-1 infection Species (MHC) human (A2, A3)

Country Viet Nam. Assay type HLA binding

Keywords subtype comparisons, computational epitope prediction, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-A11, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- ILGLNKIV is the epitope in the HXB2 reference strain sequence, and is also the most common form in CRF01.

**HXB2 Location** p24 (135–143) Author Location Gag (267–275)

**Epitope ILGLNKIVR** 

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: OS21

Species (MHC) human (A3, A11)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

Keywords vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

· After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

HXB2 Location p24 (135–145)

**Author Location** Gag (267–277)

Epitope ILGLNKIVRMY

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (B7 supertype)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

- After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the most frequently recognized of the 31 epitopes that the vaccinated volunteers responded to.
- A response to this peptide was induced in three patients after immunization with lipopeptides alone (no adjuvant) after the third and the fourth boost, and induced in two patients after immunization with lipopeptides and QS21 adjuvant after the third boost. Variant IyGLNKIVRMY was also recognized in two patients.

**HXB2 Location** p24 (136–145)

**Author Location** p24 (268–277 LAI)

Epitope LGLNKIVRMY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B62)

Keywords review

References McMichael & Walker 1994

- Predicted from larger peptide.
- Review of HIV CTL epitopes.
- Also P. Johnson, pers. comm.

**HXB2 Location** p24 (136–146)

Author Location p24 (271–281)

Epitope LGLNKIVRMYS

Immunogen HIV-1 infection Species (MHC) human (B62)

Keywords TCR usage

References Lubaki et al. 1997

- Eighty two HIV-1-specific CTL clones from 5 long-term nonprogressors were isolated and analyzed for breadth of response.
- A sustained Gag, Env and Nef response was observed, and clones were restricted by multiple HLA molecules, indicating a polyclonal response.
- A subject who was B62+ had CTL that recognized this peptide, p17 KIRLRPGGKKKYKL, and one additional unknown epitope.
- The two clones that recognized this epitope used two different  $V\beta$  genes, further demonstrating a polyclonal response.

**HXB2 Location** p24 (136–146)

Author Location p24 (136–146)

Epitope LGLNKIVRMYS

Immunogen HIV-1 infection

Species (MHC) human (B62)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location p24 (136–150)

Author Location p24 (136–150 HXB2)

**Epitope** LGLNKIVRMYSPTSI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

 $\label{lem:keywords} \textbf{Keywords} \ \ \text{supervised treatment interruptions (STI), im-}$ 

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized—the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized. A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- Responses to this peptide were detected in 16% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p24 (137–145)

Author Location p24 (272–280 LAI)

Epitope GLNKIVRMY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1501)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1501 epitope.

**HXB2 Location** p24 (137–145)

Author Location Gag (269-277 SUMA)

Epitope GLNKIVRMY

Epitope name Gag GY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1501)

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, acute/early infection, characterizing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4
   T cell counts through seven years of follow up. In contrast
   to more rapid progressors, WEAU and BORI, SUMA a broad
   response to 24 epitopes, with little immunodominance. Two
   peptides were somewhat more intensely recognized in acute
   infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

**HXB2 Location** p24 (137–145)

Author Location Gag (B con)

Epitope GLNKIVRMY

Epitope name GY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B15)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords variant cross-recognition or cross-

neutralization

References Draenert et al. 2004c

- CD8+ T-cell responses that persisted in individuals with advanced disease were studied; most of the optimal epitopes defined were recognized with intermediate to high avidity. On average 13 (range, 2-39) epitopic regions were targeted in an average of 6 proteins (range, 1-8). HAART resulted in decrease in antigen and reduction in gamma IFN EliSpot responses, suggesting active responses to autologous virus. The lack of mutations within most viral epitopes suggest that persistent CTL through late disease do not exert strong immune selection pressure, yet the Elispot assays show robust responses, suggesting to the authors that gamma IFN-based screening methods may not reveal functional CD8+ T-cell impairment in patients with AIDS.
- 1 subject recognized this epitope with intermediate functional avidity. The autologous sequence matched the B consensus.

**HXB2 Location** p24 (137–145)

Author Location p24 (272–280 LAI)

Epitope GLNKIVRMY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B62)

**Keywords** review, escape **References** Goulder *et al.* 1997a

- This paper is a review of CTL and immune evasion, but it presents a study of a shift from an HLA-A\*0201 response to SLYNTVATL, to a B62 response to GLNKIVRMY.
- As long as a strong CTL response to SLYNTVATL was evident, the epitope variants SLFNTVATL or SLYNTIATL dominated the viral population – eventually the CTL response to the index peptide became undetectable, the CTL response shifted to a focus on GLNKIVRMY, and the index peptide SLYNTVATL once again established itself as the dominant form.

**HXB2 Location** p24 (137–145)

**Author Location** p24 (SF2)

Epitope GLNKIVRMY

Immunogen HIV-1 infection

Species (MHC) human (B62)

**Keywords** subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in a HIV+ African American living in Boston this epitope did not fall within the three most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

**HXB2 Location** p24 (137–145)

Author Location p24 (267–277 SF2)

Epitope GLNKIVRMY

Immunogen HIV-1 infection

Species (MHC) human (B62)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B62+ individuals that had a CTL response to this epitope broken down by group: 0/1 group 1, 0/1 group 2, and 1/1 group 3.

**HXB2 Location** p24 (137–145)

**Author Location** p24 (137–145)

Epitope GLNKIVRMY Immunogen HIV-1 infection Species (MHC) human (B62) Keywords immunodominance References Day et al. 2001

• No immunodominant responses were detected to four B62restricted epitopes tested.

**HXB2 Location** p24 (137–145) **Author Location** p24 (137–145) Epitope GLNKIVRMY Subtype B

Immunogen HIV-1 infection Species (MHC) human (B62)

**Donor MHC** A1, A3, B8, B62, Cw3, Cw7 Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, earlyexpressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFNsecreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

**HXB2 Location** p24 (137–145) **Author Location** Gag (269–277) **Epitope** GLNKIVRMY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B62)

**Donor MHC** A2, A24, B27, B62 Assay type Chromium-release assay

Keywords TCR usage, genital and mucosal immunity

References Musey et al. 2003

• CTL clones from blood, semen, cervix and rectum samples from 12 individuals were compared. CTL clones derived from blood and mucosal samples had similar high lysis efficiency, primarily perforin dependent, and TCRbeta VDJ region sequencing revealed cases of expansion of the same clone in different compartments.

- CD8+ T cell clones directed at this epitope were derived from blood, rectum and semen.
- The TCRbeta VDJ rearrangement of the CTL clones was  $V\beta$ 22S1DJ1.2, demonstrating expansion of CTL clones in all three compartments from the same progenitor cell.

**HXB2 Location** p24 (137–145)

Author Location Gag (269-277)

Epitope GLNKIVRMY

Epitope name GY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B62)

**Donor MHC** A\*01, A\*11, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords escape, optimal epitope References Koibuchi et al. 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates
- during the chronic phase of infection. The autologous form of the GY9 matched the B clade consensus form of the epitope, GLNKIVRMY, throughout the 5 years

limited viral evolution within targeted CD8 T-cell epitopes

**HXB2 Location** p24 (137–145)

of study.

**Author Location** Gag (269–277 HXB2)

**Epitope** GLNKVRMY

Subtype B, CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (B62)

Country Viet Nam.

Assay type HLA binding

**Keywords** subtype comparisons, computational epitope

prediction, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-A11, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- GLNKVRMY is the epitope in the HXB2 reference strain sequence, and is also the most common form in CRF01.

**HXB2 Location** p24 (137–145)

Author Location p24 (C consensus)

Epitope GLNKIVRMY

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A2, B\*5802, B62, Cw4, Cw6

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in a HIV+ South African living in Durban, HLA A2/- B5802/62 Cw4/6 – this epitope did not fall within the three most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

**HXB2 Location** p24 (142–150)

**Author Location** 

Epitope VRMYSPVSI

Epitope name VI9

Immunogen

Species (MHC) human (Cw\*18)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a Cw18 epitope.

**HXB2 Location** p24 (142–150)

**Author Location** (C consensus)

Epitope VRMYSPVSI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- VRMYSPVSI is an optimal epitope.

**HXB2 Location** p24 (143–150)

Author Location p24 (273–283 IIIB)

Epitope RMYSPTSI

Immunogen HIV-1 infection

Species (MHC) human (B\*5201)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*5201 epitope.

**HXB2 Location** p24 (143–150)

**Author Location** p24 (273–283 IIIB)

**Epitope** RMYSPTSI

Epitope name SL9

Immunogen HIV-1 infection

Species (MHC) human (B52)

**Keywords** epitope processing, immunodominance, escape

References Brander et al. 1999

- Multiple natural variations in the SL9 flanking regions of the immunodominant epitope SLYNTVATL were tested and found not to adversely affect CTL recognition or prevent epitope processing, suggesting that viral escape from the HLA-A\*0201restricted CTL response against SLYNTVATL is probably not linked to variations in the flanking regions of this epitope.
- The CTL response to RMYSPTSI was used as a control.

**HXB2 Location** p24 (143–150)

Author Location p24 (273–283 IIIB)

Epitope RMYSPTSI

Immunogen HIV-1 infection

Species (MHC) human (B52)

**Keywords** responses in children, mother-to-infant transmission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- No variants of this epitope were found in a non-transmitting mother that had a CTL response to this epitope.

**HXB2 Location** p24 (143–150)

**Author Location** p24 (143–150)

Epitope RMYSPTSI

Immunogen HIV-1 infection

Species (MHC) human (B52)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (143–150)

Author Location Gag (275–282)

Epitope RMYSPTSI

Epitope name RI8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B52)

**Donor MHC** A\*02, A\*68, B\*14, B\*52, Cw\*08, Cw\*12

Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords escape, optimal epitope

References Koibuchi et al. 2005

HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.

• A form of this epitope that elicited a diminished Elispot response, RMYSPvSI, dominated the viral sequence for several years, and then reverted back to the B consensus form, • CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could RMYSPTSI.

**HXB2 Location** p24 (143–151) Author Location p24 (143–151) Epitope RMYSPVSIL Subtype C Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

- PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.
- This epitope is highly conserved across clades.

**HXB2 Location** p24 (143–151) Author Location Gag (275–283) **Epitope** RMYSPTSIL Subtype B

> Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (A2)

Immunogen vaccine

Assay type proliferation, CD8 T-cell Elispot - IFNy, Chromium-release assay

Keywords vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

• After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

**HXB2 Location** p24 (143–151) Author Location Gag (275–283 BRU) **Epitope** RMYSPTSIL Subtype B, CRF02\_AG Immunogen HIV-1 infection Species (MHC) human (A2) Country Cote D'Ivoire. Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons References Inwoley et al. 2005

- recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects
- This epitope was recognized by 1/9 CRF02 AG-infected Ivorians, and 0/9 B-infected French subjects.

**HXB2 Location** p24 (143–151)

Author Location Gag (275–283 HXB2)

**Epitope** RMYSPTSIL Subtype B, CRF01 AE Immunogen HIV-1 infection Species (MHC) human (A2)

Country Viet Nam. Assay type HLA binding

Keywords subtype comparisons, computational epitope prediction, escape, variant cross-recognition or cross-neutralization, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-A11, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- The most common CRF01\_AE variant rmyspVsil had a higher HLA-binding score than the HXB2 epitope. The rare variant, rmyspVsiW was predicted not to bind to A2.

**HXB2 Location** p24 (144–151) Author Location Gag (276–283)

**Epitope** MYSPTSIL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: OS21

Species (MHC) human (A24)

Assay type proliferation, CD8 T-cell Elispot - IFNy, Chromium-release assay

Keywords vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

• After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the most frequently recognized of the 31 epitopes that the vaccinated volunteers responded to.

HXB2 Location p24 (151-170) Author Location p24 (283–302 SF2) Epitope LDIRQGPKEPFRDYVDRFYK

Immunogen HIV-1 infection

Species (MHC) human

References McAdam et al. 1998

**HXB2 Location** p24 (155–177) **Author Location** p24 (287–309)

Epitope QGPKEPFRDYVDRFYKTLRAEQA

Immunogen vaccine

*Vector/Type:* peptide *HIV component:* p24 Gag

Species (MHC) mouse

References Nakamura et al. 1997

- Mice immunized with this synthetic peptide generated specific CTLs, a proliferative response, and antibodies.
- The amino acids shown in the epitope field were based on the numbering provided by Nakamura et al., and may not be correct.
- The CTL epitope was shown to be located in positions 291-300.

**HXB2 Location** p24 (157–178) **Author Location** p24 (290–309)

Epitope PKEPFRDYVDRFYKTLRAEQAS

Immunogen HIV-1 infection Species (MHC) human (B14) References Musey *et al.* 1997

• Cervical and peripheral blood derived CTL clones from an HIV-infected woman recognized this epitope.

**HXB2 Location** p24 (159–168) **Author Location** p24 (159–168)

Epitope EPFRDYVDRF

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Country India.

**Assay type** CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** p24 (159–168)

**Author Location** Gag (291–300)

Epitope EPFRDYVDRF

Immunogen vaccine

Vector/Type: DNA, DNA with protein boost Strain: B clade LAI HIV component: Gag,

Nef, Tat Adjuvant: IL-18

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1

References Billaut-Mulot et al. 2001

- DNA vaccinated BALB/c mice primed and boosted with the multiepitopic vaccine with IL18 showed lymphoproliferative responses 7 weeks post immunization.
- Strong but non-lasting HIV-specific CTL responses were detected by a Cr-release assay and DNA prime/DNA boost was more effective than DNA prime protein boost.
- Immunization with either the multiepitopic DNA or with the mixed DNA vaccine induced HIV-1 specific Th1 cytokines (IL-2 and IFN-gamma)
- Co-administration of IL18 increased T-cell responses but decreased anti-HIV antibody levels.

HXB2 Location p24 (159–168)

Author Location p24

Epitope EPFRDYVDRF

Epitope name E10F

Immunogen vaccine

Vector/Type: DNA HIV component: Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type Chromium-release assay

References Bojak et al. 2002b

 Codon-optimized gag gene DNA vaccines were compared to wild type by vaccination of BALB/c mice. Codon optimized DNA gave a Th1 polarized Th response, a strong antibody response that persisted from more than 20 weeks, and CTL responses, while wild-type DNA induced weak and inconsistent immune responses.

**HXB2 Location** p24 (159–168)

**Author Location** 

Epitope EPFRDYVDRF Epitope name E10F Immunogen vaccine

Vector/Type: DNA, virus-like particle (VLP), polyepitope HIV component: Gag, p24 Gag,

**Species (MHC)** mouse (H-2L<sup>d</sup>)

Assay type Cytokine production, Chromium-release assay

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance

References Wild et al. 2004

- A codon optimized gag DNA vaccine was compared to a myristylation defective gag and p24 alone, both of which lack signals for secretion from transfected cells. Gag-derived immunogens that were secreted as VLPs and those that remained intracellular (p24) each produced strong CTL responses, and neither the size of antigen nor cellular trafficking and localization significantly influenced the strength of humoral and cellular immune activation. The formation and release of VLPs was not essential for eliciting strong CTL. BALB/c mice were given the DNA vaccine by i.m. administration of plasmid DNA for the prime and boost.
- Minigenes were made incorporating just 1 epitope, minitopes, carrying 1 of 3 murine class I epitopes linked to the Ad2-E3 protein-derived signal peptide to allow access of the epitope to the ER. Weak induction of cellular immune responses was

observed, in contrast to the complex polyprotein. The E10F minigene did not produce a detectable CTL response.

**HXB2 Location** p24 (159–178) **Author Location** Gag (96ZM651.8)

Epitope EPFRDYVDRFFKTLRAEQAT

Immunogen

Species (MHC) human (B\*440301)

Keywords subtype comparisons, immunodominance

References Novitsky et al. 2001

- This study is provides a survey of CTL responses and full length HIV-1 genome sequences from a C subtype infected Botswanan cohort.
- 16 of 46 (34.8%) had CTL responses to one or more peptides within the second immunodominant region region of Gag (peptides SILDIKQGPKEPFRDYVDRF, EPFRDYVDRFFK-TLRAEQAT, and FKTLRAEQATQEVKNWMTDT) with ELISPOT results median and range 500 (100 to 1,250) SFC/10<sup>6</sup> PBMC
- 3 of 6 (50%) carriers of HLA-B\*44031 showed CTL responses to the peptide EPFRDYVDRFFKTLRAEQAT.

**HXB2 Location** p24 (159–178)

Author Location Gag (291–310)

Epitope EPFRDYVDRFFKTLRAEQAT

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons

References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 8 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** p24 (160–169)

Author Location p24

Epitope PFRDYVDRFF

Epitope name PF-10

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining,

Characian allere and eyeoki

Chromium-release assay

**Keywords** subtype comparisons, epitope processing, immunodominance, cross-presentation by differ-

ent HLA

References Masemola et al. 2004b

 Highly targeted regions in Gag for CD8+ T-cells were defined for individuals with C clade infections in South Africa. 9 specific epitopes within the most reactive regions were characterized. This is 1 of 5 novel epitopes that were found among subtype C HIV-1 from African patients who hadn't previously been identified in B clade infections. Some epitopes were shown to be promiscuous, presented by multiple class I restricting alleles. The HLA restricting element for this optimal epitope was not determined due to limited material.

**HXB2 Location** p24 (161–169)

**Author Location** 

Epitope FRDYVDRFF

Immunogen

Species (MHC) human (Cw\*18)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an Cw18 epitope.

HXB2 Location p24 (161-169)

**Author Location** (C consensus)

Epitope FRDYVDRFF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (161–169)

**Author Location** (C consensus)

Epitope FRDYVDRFF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- FRDYVDRFF is an optimal epitope.

**HXB2 Location** p24 (161–170)

**Author Location** p24 (subtype B, D)

Epitope FRDYVDRFYK

Subtype B, D

Immunogen HIV-1 infection

Species (MHC) human (B\*1801)

References Ogg et al. 1998a

DRFY.

**HXB2 Location** p24 (161–170)

**Author Location** p24 (subtype B, D)

Epitope FRDYVDRFYK

Subtype B, D

Immunogen HIV-1 infection

Species (MHC) human (B\*1801)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1801 epitope.

**HXB2 Location** p24 (161–170)

**Author Location** p24 (161–170 HXB2)

Epitope FRDYVDRFYK

Epitope name FK10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1801)

**Donor MHC** A\*0201, A\*2501, B\*1801. B\*5101.

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

• T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.

HXB2 Location p24 (161-170)

Author Location p24 (161–170)

Epitope FRDYVDRFYK

Immunogen HIV-1 infection

Species (MHC) human (B18)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (161–170)

Author Location p24 (293–302)

Epitope FRDYVDRFYK

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B18)

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2001a

- Variants FRDYVDRF(Y/F)K are specific for the B,D/A,C clades.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

- Noted in Brander 1999, this database, to be B\*1801, FRDYV- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
  - 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
  - Among HLA-B18 women, 3/4 HEPS and 1/9 HIV-1 infected women recognized this epitope, likelihood ratio 5.3, p value 0.04, and HEPS women tended to respond to FRDYV-DRFY/FK, while infected women tended to respond to YPLT-FGWCY/F.
  - The dominant response to this HLA allele was to this epitope for all 3/4 HEPS cases and for the single HIV-1 infected women that responded to this epitope.
  - Four epitopes were considered to be "resistant epitopes", as they were preferentially reactive in HEPS women and so may confer resistance, and these were found in three different proteins: A2 ILK(D/E)PVHGV in RT, A\*6802 DTVLEDINL in Protease, B14 DLNM/TLN(I/V)V in p24 and B18 FRDYV-DRF(Y/F)K also in p24.
  - Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.

HXB2 Location p24 (161-170)

**Author Location** p24

Epitope FRDYVDRFYK

Subtype B, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade

HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (B18)

Keywords subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

**References** Hanke & McMichael 2000: Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location p24 (161-170)

**Author Location** 

**Epitope** FRDYVDRFFK **Immunogen** HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls, ML1732.

**HXB2 Location** p24 (161–174)

**Author Location** p24 (161–174 HXB2)

Epitope FRDYVDRFYKTLRA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

 $\textbf{Keywords} \ \ \text{supervised treatment interruptions (STI), im-}$ 

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- Responses to this peptide were detected in 16% of the study subjects, and it was one of the 25 most frequently recognized peptides.

HXB2 Location p24 (161-175)

**Author Location** p24 (161–170)

Epitope FRDYVDRFYKTLRAE

Subtype A, D

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*0101, A\*7401, B\*5801

Country Uganda.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References Barugahare et al. 2005

- T-cell immune responses were examined in HIV-1 infected Ugandans. Similar levels of cross-clade responses for Gag, Env and Nef were observed. Higher frequencies of responses in conserved regions were found for Gag within a single clade, while areas with higher viral sequence variation had fewer detectable responses. Differential interclade sequence homologies between the Gag regions did not affect the respective level of cross-clade recognition.
- The sequence contains a known epitope (FRDYVDRFYKTL), but the subject recognizing it does not carry HLAs of the previously-defined restriction. The isolated viral sequence was frdyvdrfykVlrae, from the patient that could recognize the peptide.

HXB2 Location p24 (161–180)

Author Location p24 (293–312 SF2)

Epitope FRDYVDRFYKTLRAEQASQD

Immunogen HIV-1 infection Species (MHC) human (B71)

References McAdam et al. 1998

**HXB2 Location** p24 (161–180)

**Author Location** p24 (293–312 SF2)

Epitope FRDYVDRFYKTLRAEQASQD

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A2, A3, B8, B62.

**HXB2 Location** p24 (161–180)

Author Location p24 (293–312 SF2)

Epitope FRDYVDRFYKTLRAEQASQD

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

• CTL expanded *ex vivo* were later infused into HIV-1 infected patients.

**HXB2 Location** p24 (162–172)

**Author Location** p24 (296–306 subtype A)

Epitope RDYVDRFFKTL

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

**Keywords** subtype comparisons

References Dorrell et al. 1999

 CTL responses in three individuals with non-clade B infections were studied, 2 with subtype A infections, 1 with subtype C – their infections all originated in East Africa.

- This epitope is similar to the A24 DYVDRYFKT epitope found for B subtype, but CTL from this A subtype infection required the additional Arg – the B clade sequence change from F to Y diminished CTL reactivity.
- C. Brander notes that this is an A\*2402 epitope in the 1999 database.

**HXB2 Location** p24 (162–172)

Author Location p24 (296–306 subtype A)

Epitope RDYVDRFFKTL

Subtype A

Immunogen HIV-1 infection

**Species (MHC)** human (A\*2402) **Keywords** optimal epitope

**References** Frahm et al. 2007

• C. Brander notes this is an A\*2402 epitope.

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**HXB2 Location** p24 (162–172) **Author Location** p24 (296–306)

Author Location p24 (296–306)

Epitope RDYVDRFFKTL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A24)

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-A24 women, 0/4 HEPS and 6/10 HIV-1 infected women recognized this epitope, likelihood ratio 7.2, p value 0.03, and (R)YL(R/K)DQQLL tended to be reactive in HEPS and infected women, RDYVDRFFKTL in infected women only.
- The dominant response to this HLA allele was to this epitope in all of the 6/10 HIV-1 infected women.
- Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.
- Subject ML 1707 started with a CTL response to A\*6802 DTVLEDINL prior to seroconversion, and switched to A\*6802 ETAYFILKL and A24 RDYVDRFFKTL post-seroconversion.

**HXB2 Location** p24 (162–172)

Author Location p24 (293–312 LAI)

 ${\bf Epitope} \ \ {\tt RDYVDRFYKTL}$ 

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*4402)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*4402 epitope.

**HXB2 Location** p24 (162–172)

**Author Location** p24 (162–172)

Epitope RDYVDRFYKTL

Immunogen HIV-1 infection

Species (MHC) human (B44)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (162–172)

**Author Location** p24 (162–172)

Epitope RDYVDRFYKTL

Immunogen HIV-1 infection

Species (MHC) human (B44)

References Day et al. 2001

HXB2 Location p24 (162–172)

Author Location p24

Epitope RDYVDRFYKTL

Subtype B, D

Immunogen HIV-1 infection, vaccine

*Vector/Type:* DNA prime with modified vaccinia Ankara (MVA) boost *Strain:* A clade

HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (B44)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses
  to peptide pools were detected using intracellular cytokine
  staining and IFNgamma Elispot assays after vaccination of 5
  macaques. The response to the Mamu A\*01 SIV p27 epitope
  p11C (CTPYDINQM), included in the polyepitope region, was
  not immunodominant in the Mamu A\*01 vaccinated macaques,
  possibly because of processing limitations in context of the
  artificial polyepitope string Wee et al. [2002].

**HXB2 Location** p24 (162–172)

Author Location Gag (B con)

Epitope RDYVDRFYKTL

Epitope name RL11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B44)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords variant cross-recognition or cross-

neutralization

References Draenert et al. 2004c

- CD8+ T-cell responses that persisted in individuals with advanced disease were studied; most of the optimal epitopes defined were recognized with intermediate to high avidity. On average 13 (range, 2-39) epitopic regions were targeted in an average of 6 proteins (range, 1-8). HAART resulted in decrease in antigen and reduction in gamma IFN EliSpot responses, suggesting active responses to autologous virus. The lack of mutations within most viral epitopes suggest that persistent CTL through late disease do not exert strong immune selection pressure, yet the Elispot assays show robust responses, suggesting to the authors that gamma IFN-based screening methods may not reveal functional CD8+ T-cell impairment in patients with AIDS.
- 1 subject recognized this epitope with low functional avidity. The autologous sequence matched the B consensus.

**HXB2 Location** p24 (162–172)

Author Location p24 (293–312 LAI)

Epitope RDYVDRFYKTL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B44, A26, B70)

References Ogg et al. 1998a

**HXB2 Location** p24 (163–171)

Author Location Gag (295–303 SUMA)

Epitope DYVDRFYKT

**Epitope name** Gag DT9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

Keywords dynamics, acute/early infection, characteriz-

ing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4 T cell counts through seven years of follow up. In contrast to more rapid progressors, WEAU and BORI, SUMA a broad response to 24 epitopes, with little immunodominance. Two peptides were somewhat more intensely recognized in acute infection, but this response leveled out early on.

• Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

**HXB2 Location** p24 (163–172)

**Author Location** p24 (163–172)

Epitope DYVDRFYKTL

Immunogen HIV-1 infection

Species (MHC) human (A24)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (163–173)

**Author Location** Gag (297–307 SF2)

Epitope DYVDRFYKTLR

Subtype B

Immunogen HIV-1 infection, computer prediction

Species (MHC) human (A\*3303)

Assay type Chromium-release assay

Keywords binding affinity, computational epitope predic-

tion

References Hossain et al. 2003

- HLA-A\*3303 is a common HLA allele in east and southeast Asia. Pol, Gag and Nef SF2 proteins were scanned for potential A\*3303 epitopes. 99 potential epitopes were synthesized, and 52/99 bound to A\*3303. Six of these served as peptide-targets for lysis by PBMC from infected individual, and clones derived from 4 of these 6 could lyse HIV-vaccinia infected target cells, indicating proper processing.
- This epitope is one of the 2/6 peptides that could induce CTL responses in the PBMC of infected individuals, but was not properly processed in a vaccinia-HIV infected target cell.

**HXB2 Location** p24 (164–172)

**Author Location** Gag (296–304)

Epitope YVDRFYKTL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0207)

Donor MHC A\*0207

**Keywords** subtype comparisons

References Currier et al. 2002a

- Cross-reactive responses were found in PBMC isolated from individuals infected with either B or CRF01\_AE clade viruses, as determined by Elispot assays of target cells expressing recombinant vaccinia viruses expressing HIV-1 gag, env, nef and pol from many clades.
- The Thai subject VAIP-4 demonstrated broad CTL cross-reactivity towards gag constructs derived from subtypes A, B, C, D, F, G, H, and CRF-01\_AE. Sequence alignments of this epitope showed conservation for clades B and D, and Y->F substitutions at position 6 for subtypes A, C, CDR01-AE, F, G, and H. YVDRFYKTL and the variant epitope YVDRFFKTL are recognized equally well.

**HXB2 Location** p24 (164–172)

Author Location p24 (164–172)

Epitope YVDRFYKTL Immunogen HIV-1 infection Species (MHC) human (A\*0207) Keywords optimal epitope References Frahm *et al.* 2007

**HXB2 Location** p24 (164–172)

Author Location p24 (298–306 subtype A)

Epitope YVDRFFKTL

Subtype A

Immunogen HIV-1 infection Species (MHC) human (A26, B70)

**Keywords** subtype comparisons **References** Dorrell *et al.* 1999

- CTL responses in three individuals with non-clade B infections were studied, 2 with subtype A infections, 1 with subtype C – their infections all originated in East Africa.
- This CTL epitope is conserved in A and C subtype, and B clade sequences tend to have a change from F to Y, YVDRFYKTL – both variants showed strong CTL reactivity.
- CTL reacted with targets presenting either in the context A26 or B70 the epitope has the HLA-26 motif of Val at position 2 and Leu at the carboxy terminus, and the B70 anchor residue motif is unknown.

**HXB2 Location** p24 (164–172)

Author Location Gag (298-306 subtype A)

Epitope YVDRFFKTL

Subtype A

Immunogen HIV-1 infection, in vitro stimulation or selec-

Species (MHC) human (A26, B70)

 $\textbf{Keywords} \ \ \text{subtype comparisons}$ 

References Dorrell et al. 2001

 In vitro restimulation of CTL specific for dominant epitopes from infected individuals is possible using recombinant modified vaccinia virus Ankara (MVA) carrying A or D subtype HIV-1 Gag proteins.

**HXB2 Location** p24 (164–172)

**Author Location** p24

Epitope YVDRFFKTL

Epitope name YL-9

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*1503)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining,

Chromium-release assay

**Keywords** subtype comparisons, epitope processing, immunodominance, cross-presentation by differ-

ent HLA

References Masemola et al. 2004b

 Highly targeted regions in Gag for CD8+ T-cells were defined for individuals with C clade infections in South Africa. 9 specific epitopes within the most reactive regions were characterized.  YVDRFFKTL was presented by B\*15, which is more common in Zulus than Caucasians (0.153 versus 0.079). This epitope had previously identified in B clade infections.

**HXB2 Location** p24 (164–172)

**Author Location** 

Epitope YVDRFFKTL

Immunogen

Species (MHC) human (B\*1503)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B\*1503 epitope.

HXB2 Location p24 (164-172)

**Author Location** Gag (296–304 96ZM651.8)

Epitope YVDRFFKRL

Immunogen

Species (MHC) human (B\*1510, B70)

**Keywords** subtype comparisons

References Novitsky et al. 2001

- This study provides a survey of CTL responses and full length HIV-1 genome sequences from a C subtype infected Botswanan cohort
- 4 subjects who responsed to the CTL epitope YVDRFFKTL all were HLA-B\*1510 and also shared HLA-Cw03, suggesting linkage disequilibrium.
- An HIV-1 B variant of the epitope YVDRFYKTL has been described, and was recognized by CTL from an HIV-1 subtype A-infected patient, and the HLA restriction of the epitope was suggested to be A26 or B70 – HLA-B\*1510 is equivalent to the serological specificity HLA B70.

HXB2 Location p24 (164-172)

**Author Location** p24 (164–172)

Epitope YVDRFYKTL

Immunogen HIV-1 infection

Species (MHC) human (B70)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (164–172)

**Author Location** p24 (164–172)

Epitope YVDRFFKTL

Immunogen

Species (MHC) human (Cw\*0303)

**Keywords** optimal epitope

References Frahm et al. 2007

**HXB2 Location** p24 (164–172)

**Author Location** p24 (164–172)

Epitope YVDRFFKTL

Immunogen

Species (MHC) human (Cw\*0304)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location p24 (164–172)

Author Location (C consensus)

 ${\bf Epitope}\ \ {\tt YVDRFFKTL}$ 

Epitope name YL9

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0304)

**Donor MHC** A\*3402, B\*0801, B\*4403, Cw\*0304,

Cw\*0401

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was one of two used to illustrate how specific epitopes were characterized with regard to defining the optimal epitope and the HLA restricting element. HLA allelic associations in the population with peptide recognition was highly predictive of the epitope within the 15 mer.

**HXB2 Location** p24 (164–172)

**Author Location** (C consensus)

**Epitope** YVDRFFKTL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0304)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the T8 residue of YVDRFFKTL are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** p24 (165–178)

Author Location p24 (165–177 HXB2)

**Epitope VDRFYKTLRAEQAS** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assav type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

 Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.

- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 19% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p24 (166–174)

**Author Location** (C consensus)

Epitope DRFFKTLRA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1401)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- DRFFKTLRA is an optimal epitope.

**HXB2 Location** p24 (166–174)

Author Location p24 (298–306 LAI)

Epitope DRFYKTLRA

**Subtype** B

Immunogen HIV-1 infection

Species (MHC) human (B\*1402)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1402 epitope.

**HXB2 Location** p24 (166–174)

**Author Location** (167–175)

Epitope DRFFKTLRA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1402)

Assay type Other

**Keywords** HLA associated polymorphism

**References** Boutwell & Essex 2007

 All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.

 DRFFKTLRA was a previously defined B\*1402 presented epitope that encompassed a B\*14/B\*1401 associated polymorphism, DRFFkTLRA,in the fifth position. This epitope is embedded in a previously determined CTL immunodominant region.

HXB2 Location p24 (166-174)

Author Location p24

Epitope DRFFKTLRA

Epitope name DA-9

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*1403)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ, Intracellular cytokine staining,

Chromium-release assay

**Keywords** subtype comparisons, epitope processing, immunodominance, cross-presentation by differ-

ent HLA

References Masemola et al. 2004b

- Highly targeted regions in Gag for CD8+ T-cells were defined for individuals with C clade infections in South Africa. 9 specific epitopes within the most reactive regions were characterized.
- DRFFKTLRA was presented by B\*14, which is more common in Zulus than Caucasians (0.066 versus 0.038). This epitope had previously identified in B clade infections.

**HXB2 Location** p24 (166–174)

Species (MHC) human (B14)

Author Location p24 (298–306 IIIB)

**Epitope** DRFYKTLRA **Immunogen** HIV-1 infection

**Keywords** responses in children, mother-to-infant transmission

References Wilson et al. 1996

- Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.
- DRFYKILRA, a naturally occurring variant, was found in mother, and is recognized although less reactive.
- DQFYKTLRA, a naturally occurring variant, was found in infant and is not recognized.

**HXB2 Location** p24 (166–174)

Author Location p24 (298–306 IIIB)

Epitope DRFYKTLRA

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Cao et al. 1997a

- The consensus peptide for clades B and D is DRFYKTLRA.
- The consensus peptide for clades A and C is DRFFKTLRA and it is equally reactive.

HXB2 Location p24 (166–174)

Author Location p24 (298–306 HXB2)

**Epitope** DRFYKTLRA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords kinetics

References Yang et al. 1997b

- A chimeric universal T cell receptor was created by linking CD4 or an HIV-specific anti-gp41 Ig sequence to the signaling domain of the T cell receptor chain ζ, and transducing into CD8+ cells.
- The response using universal-receptor-bearing CD8+ cells to lyse infected cells *in vitro* was comparable to the natural occurring responses of CTL-clones from HIV+ individuals in terms of kinetics and efficiency.
- A CTL clone specific for this epitope was used for the comparison.

HXB2 Location p24 (166-174)

**Author Location** p24

Epitope DRFWKTLRA

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B14)

**Keywords** subtype comparisons, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades such cross-reactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The D subtype consensus is identical to the B clade epitope.
- The A subtype consensus is drFfKtLRA.

HXB2 Location p24 (166–174)

Author Location p24 (298–306 LAI)

Epitope DRFYKTLRA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Harrer et al. 1996b

**HXB2 Location** p24 (166–174)

Author Location p24 (298–306)

Epitope DRFYKTLRA

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Yang et al. 1996

- CD4+ cell lines acutely infected with HIV were studied to determine their susceptibility to lysis by CTL.
- Clones specific for RT lysed HIV-1 infected cells at lower levels than Env or Gag specific clones.
- The distinction was thought to be due to lower expression of RT relative to Env and Gag.
- CTL can lyse infected cells early after infection, possibly prior to viral production.

HXB2 Location p24 (166–174)

Author Location p24 (298–306)

Epitope DRFYKTLRA Immunogen HIV-1 infection Species (MHC) human (B14)

Assay type CTL suppression of replication

References Yang et al. 1997a

- CTL inhibit HIV-1 replication at effector cell concentrations comparable to those found *in vivo*.
- CTL produced HIV-1-suppressive soluble factors MIP-1α, MIP-1β, RANTES, after antigen-specific activation.
- CTL suppress HIV replication more efficiently in HLAmatched cells.

**HXB2 Location** p24 (166–174) **Author Location** p24 (298–306)

Epitope DRFYKTLRA

Immunogen in vitro stimulation or selection

Species (MHC) human (B14)

Keywords dendritic cells

References Zarling *et al.* 1999

- This study compares the ability of macrophages and dendritic cells to stimulate primary responses in CD8+ lymphocytes isolated from HLA-appropriate HIV-uninfected donors using peptide-pulsed APC – the dendritic cells performed better as APC for the stimulation of primary responses.
- Strong CTL responses were elicited by the epitopes DRFYK-TLRA and GEIYKRWII when presented by either immature or mature dendritic cells – macrophages were not able to prime a CTL response against DRFYKTLRA.
- A weak response to KLTPLCVSL was stimulated using macrophages as the APC.
- No detectable response was observed for the following previously-defined HIV epitopes: KIRLRPGGK, ILKEPVHGV, IRLRPGGK, GPKVKQWPL.

**HXB2 Location** p24 (166–174)

Author Location p24

Epitope DRFYKLTRA

Immunogen

Species (MHC) human (B14)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied – these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive, and the B35 allele seems to be protective.
- HIV-2 sequence: DRFYKSLRA is cross-reactive, Harrer *et al.* [1993]

**HXB2 Location** p24 (166–174)

Author Location p24 (298–306 IIIB)

Epitope DRFYKTLRA
Immunogen HIV-1 infection
Species (MHC) human (B14)

Keywords responses in children, mother-to-infant trans-

mission, escape

References Wilson et al. 1999a

 This study describes maternal CTL responses in the context of mother-to-infant transmission.

- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- DRFYKILRA and DQFYKTLRA were escape mutants.

**HXB2 Location** p24 (166–174)

Author Location p24 (SF2)

Epitope DRFYKTLRA

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in 2/5 HIV+ individuals who were HLA B14 living in Boston – this epitope did not fall within the three most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

**HXB2 Location** p24 (166–174)

Author Location p24 (SF2)

Epitope DRFYKTLRA

Epitope name DA9

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords acute/early infection

References Goulder et al. 2001a

- Data from patient AC13 suggest a role for this epitope in initial control of viremia in acute infection, as it is one of several subdominant CTL epitopes recognized during the initial decline in viremia.
- A CTL response to SL9, SLYNTVATL, was not evident until 18 months post-presentation.

**HXB2 Location** p24 (166–174)

**Author Location** p24 (166–174)

Epitope DRFYKTLRA

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (166–174)

Author Location p24 (298–306 SF2)

Epitope DRFYKTLRA

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B14+ individuals that had a CTL response to this epitope broken down by group: 3/3 group 1, 1/2 group 2, and 0/0 group 3.

HXB2 Location p24 (166–174)

**Author Location** p24 (298–306)

Epitope DRFFKTLRA

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (B14)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- Variants DRF(F/W)KTLRA are specific for clades A/B.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B14 women, 0/4 HEPS and 6/7 HIV-1 infected women recognized this epitope, likelihood ratio 14.4, p value 0.004 and HEPS women tended to respond to DLNMML-NIV/DLNTMLNVV, while infected women tended to respond to DRF(F/W)KTLRA.
- The dominant response to this HLA allele was to this epitope for all of the 6/7 HIV-1 infected women.
- Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.

HXB2 Location p24 (166-174)

Author Location p24 (SF2)

Epitope DRFYKTLRA

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Altfeld et al. 2000

• This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.

HXB2 Location p24 (166–174)

**Author Location** p24

**Epitope** DRFYKTLRA

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords epitope processing

References Cao et al. 2002

- AC13 is a B14 restricted CTL clone that recognizes DRFYK-TLRA.
- CTL could be activated by a fusion protein of an HIV protein and anthrax lethal factor (LFn-HIV) that promotes antigen presenting cell uptake of exogenous protein and allows processing through the MHC class I pathway. This strategy for CTL detection could allow antigen presentation without generation of cells by the standard methods of using live viral vectors carrying a protein, or by loading the cells with peptides and by-passing processing.

**HXB2 Location** p24 (166–174)

**Author Location** p24

Epitope DRFWKTLRA

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

**HXB2 Location** p24 (166–174)

Author Location p24

Epitope DRFYKTLRA

Subtype B, D

Immunogen HIV-1 infection, vaccine

*Vector/Type:* DNA prime with modified vaccinia Ankara (MVA) boost *Strain:* A clade *HIV component:* p17 Gag, p24 Gag

Species (MHC) human (B14)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

• The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA prime-boost vaccination protocol using the HIVA antigen will be used

in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].

• Killing efficiency of CTLs and their ability to suppress viral replication is shown to depend more on epitope specificity than

• Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

**HXB2 Location** p24 (166–174) **Author Location** p24 (166–174)

Epitope DRFYKTLRA

Subtype B

**Immunogen** HIV-1 infection **Species** (MHC) human (B14)

**Donor MHC** A1, A3, B7, B14, Cw\*0702, Cw\*0802; A1,

A1, B8, B14, Cw7, Cw8

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

 $\textbf{Keywords} \ \ \text{acute/early infection, early-expressed proteins}$ 

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- This epitope was recognized in two subjects early in infection, presented by B14 in each case.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

**HXB2 Location** p24 (166–174)

Author Location p24 (166–174)

Epitope DRFYKTLRA

Epitope name Gag/p24-DA9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B14)

Assay type Chromium-release assay

Keywords binding affinity, TCR usage, characterizing

CD8+ T cells

References Yang et al. 2003b

- Killing efficiency of CTLs and their ability to suppress viral replication is shown to depend more on epitope specificity than to antigenic avidity. Different clones recognizing the same epitope had similar killing efficiency despite their variation in avidity. Nef specific CTL clones tended to be most inhibitory, followed by Gag, then by RT specific clones, regardless of avidity.
- 1/14 CTL T-cell clones tested were specific for Gag/p24-DA9. Under conditions of excess peptide (100ug/ml), there was no difference in their lytic potential; all possessed similar effector capacity. Avidity was measured as the sensitizing dose of peptide required for 50% of maximal killing (SD50), which varied from 20 pg/ml to 100 ng/ml, over four orders of magnitude for all 14 epitopes. The SD50 value for Gag/p24-DA9 was 100,000 pg/ml, it had the lowest avidity of the 14 tested.

HXB2 Location p24 (166-174)

**Author Location** (C consensus)

Epitope DRFFKTLRA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B14)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (166–174)

Author Location (B consensus)

Epitope DRFYKTLRA

Epitope name DA9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Donor MHC** A28, A29, B14, B44, Cw8; A25, A32, B08,

B14, Cw7, Cw8; A03, B14, B60, Cw3, Cw7

Country United States.

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

 Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1-specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.

 3/9 individuals recognized this epitope, presented by HLA-B14.

HXB2 Location p24 (166–174) Author Location Gag (298–306) Epitope DRFYKTLRA Subtype B

Immunogen HIV-1 infection Species (MHC) human (B14)

Donor MHC A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** p24 (166–174)

Author Location Gag (298–306)

Epitope DRFYKTLRA

Epitope name DA9 Subtype B

Immunogen HIV-1 infection Species (MHC) human (B14)

**Donor MHC** A\*02, A\*68, B\*14, B\*52, Cw\*08, Cw\*12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** escape, optimal epitope **References** Koibuchi *et al.* 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes
- A form of this epitope that elicited a diminished Elispot response, DRFYrTLRA, dominated the viral sequence for several years, and then reverted back to the B consensus form, DRFYKTLRA.

**HXB2 Location** p24 (166–174)

Author Location p24

Epitope DRFYKTLRA

during the chronic phase of infection.

Epitope name DA9

Immunogen

Species (MHC) (B14)

**Keywords** review, immunodominance, escape, acute/early infection, early-expressed proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

This review discusses the importance of 3 factors that impact
the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA
class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides
listed as immunodominant in acute HIV-1 infection.

 $\begin{array}{c} \textbf{HXB2 Location} & p24 \ (166-174) \\ \textbf{Author Location} & p24 \ (subtype \ B) \\ \textbf{Epitope} & \mathsf{DRFYKTLRA} \end{array}$ 

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B14, B\*1402)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among B and D clade viruses.
- The Clade A version of the epitope, DRFFKLTRA, was preferentially recognized by CTL.
- This epitope was recognized by two different exposed and uninfected prostitutes.

HXB2 Location p24 (166-175)

**Author Location** p24 (298–306 HX10)

Epitope DRFYKTLRAE

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords immunodominance

References Wagner et al. 1999

- The immunodominant CTL response in a long-term survivor was to this highly conserved and functionally relevant epitope.
- By testing mutations in an HXB2 background, it was found that all mutations within the epitope that abrogated CTL recognition also abolished viral infectivity.
- The epitope in this study overlaps the major homology region for which highly conserved residues exist in all known lentiand onco-viruses and yeast transposons.
- Patient was part of the study in Harrer et al. [1996a]

**HXB2 Location** p24 (166–175)

Author Location Gag (298–307)

Epitope DRFYKTRAE

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Donor MHC** A24, A33, B14, B27

Assay type Chromium-release assay

Keywords TCR usage, genital and mucosal immunity

References Musey et al. 2003

- CTL clones from blood, semen, cervix and rectum samples from 12 individuals were compared. CTL clones derived from blood and mucosal samples had similar high lysis efficiency, primarily perforin dependent, and TCRbeta VDJ region sequencing revealed cases of expansion of the same clone in different compartments.
- CD8+ T cell clones directed at this epitope were derived from blood and cervix.

**HXB2 Location** p24 (166–176)

Author Location Gag (295–305 BORI)

Epitope DRFYKTLRAEQ

Epitope name Gag DQ11

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*1402)

Donor MHC A\*2902, B\*1402, Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, immunodominance, acute/early infection, characterizing CD8+ T cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient BORI had high viral loads and rapid CD4 decline.
   BORI mounted 8 detected responses. BORI did not control viral replication well, and escape mutations occurred early and in most epitopes, 6/7 that were precisely identified.
- DRFYKTLRAEQ didn't vary. There was no response in acute infection to this epitope, but the response was detectable by early infection.

**HXB2 Location** p24 (169–185)

Author Location p24 (169–184 HXB2)

Epitope YKTLRAEQASQDVKNWN

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

 Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.

- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- Responses to this peptide were detected in 17% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p24 (169–188)

**Author Location** Gag (301–320)

Epitope YKTLRAEQASQEVKNWMTET

Immunogen HIV-1 infection Species (MHC) human (B57) Donor MHC A1, A66, B52, B57 Assay type Chromium-release assay

**Keywords** TCR usage, genital and mucosal immunity

References Musey et al. 2003

- CTL clones from blood, semen, cervix and rectum samples from 12 individuals were compared. CTL clones derived from blood and mucosal samples had similar high lysis efficiency, primarily perforin dependent, and TCRbeta VDJ region sequencing revealed cases of expansion of the same clone in different compartments.
- CD8+ T cell clones directed at this epitope were derived from blood and rectum.

**HXB2 Location** p24 (169–188)

Author Location Gag (301–320)

Epitope FKTLRAEQATQDVKNWMTDT

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 8 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location p24 (171-180)

Author Location p24

Epitope TLRAEQATQD

Epitope name TD-10

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0304)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, Chromium-release assay

**Keywords** subtype comparisons, epitope processing, immunodominance, cross-presentation by differ-

ent HLA

References Masemola et al. 2004b

- Highly targeted regions in Gag for CD8+ T-cells were defined for individuals with C clade infections in South Africa. 9 specific epitopes within the most reactive regions were characterized. This is 1 of 5 novel epitopes that were found among subtype C HIV-1 from African patients who hadn't previously been identified in B clade infections. Some epitopes were shown to be promiscuous, presented by multiple class I restricting alleles.
- TLRAEQATQD was presented by Cw\*03 and newly identified in this study; Cw\*03 is more common in Zulus than Caucasians (0.157 versus 0.101).

**HXB2 Location** p24 (173–181)

Author Location Gag (305–313 SUMA)

Epitope RAEQASQEV

Epitope name Gag RV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0802)

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

Keywords dynamics, acute/early infection, characteriz-

ing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4
   T cell counts through seven years of follow up. In contrast
   to more rapid progressors, WEAU and BORI, SUMA a broad
   response to 24 epitopes, with little immunodominance. Two
   peptides were somewhat more intensely recognized in acute
   infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

HXB2 Location p24 (173-181)

Author Location p24 (305–313)

Epitope RAEQASQEV

Immunogen HIV-1 infection

Species (MHC) human (Cw8)

References Johnson et al. 1991

- Originally reported as HLA-B14 restricted, but subsequently found not to be presented by cells transfected with B14.
- Thought to be HLA-Cw8 restricted (C. Brander and B. Walker)

**HXB2 Location** p24 (173–181)

**Author Location** p24

**Epitope** RAEQASQEV

Immunogen HIV-1 exposed seronegative

Species (MHC) human (Cw8)

**Keywords** subtype comparisons, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades such cross-reactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A subtype consensus is RAeQAtQEV.
- The D subtype consensus is RAEQsQdV.
- Thought to be HLA-Cw8 restricted, not B14 as originally reported (C. Brander, B. Walker, and S. Rowland-Jones, personal communication)

HXB2 Location p24 (173-181)

Author Location p24 (305–313)

**Epitope** RAEQASQEV

Immunogen HIV-1 infection

Species (MHC) human (Cw8)

References Price et al. 1995

- Study of cytokines released by HIV-1 specific activated CTL.
- Thought to be HLA-Cw8 restricted, not B14 as originally reported (C. Brander, B. Walker, and S. Rowland-Jones, personal communication)

**HXB2 Location** p24 (173–181)

**Author Location** p24 (305–313)

Epitope RAEQASQEV

Immunogen HIV-1 infection

Species (MHC) human (Cw8)

References Lubaki et al. 1997

- Eighty two HIV-1-specific CTL clones from 5 long-term non-progressors were isolated and analyzed for breadth of response.
- A sustained Gag, Env and Nef response was observed, and clones were restricted by multiple HLA epitopes, indicating a polyclonal response.
- Despite this being a well defined conserved epitope, and thought to be presented by B14, none of the 11 gag-specific clones from a B-14 positive subject could recognize either it or p24 PODLNTMLN.
- Thought to be HLA-Cw8 restricted, not B14 as originally reported (C. Brander, B. Walker, and S. Rowland-Jones, personal communication)

**HXB2 Location** p24 (173–181)

**Author Location** p24 (305–313)

Epitope RAEQASQEV

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (Cw8)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers. HXB2 Location p24 (173–181)

Author Location Gag (305–313)

Epitope RAEQASQEV

Epitope name RV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw8)

Donor MHC A\*02, A\*68, B\*14, B\*52, Cw\*08, Cw\*12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** escape, optimal epitope **References** Koibuchi *et al.* 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- Elispot responses to the consensus form of this epitope, RAE-QASQEV, were much more intense than to the most common variants of the epitope found over time in this individual, RAE-QAStEV and RAEQASQdV. There was a diminished response to RAEQAStEV and no response to RAEQASQdV, so these appear to be escape variants. The strong response to the consensus form persisted, despite the fact it was not observed among the autologous sequences until it surfaced as a minor variant (5/13 sequences) after 6 years of chronic infection.

**HXB2 Location** p24 (173–181)

**Author Location** 

**Epitope** RAEQASQEV **Immunogen** HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls ML1792.

HXB2 Location p24 (174–184) Author Location p24 (306–316 LAI) Epitope AEQASQDVKNW Subtype B

Immunogen

230

Species (MHC) human (B\*4402)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*4402 epitope.

HXB2 Location p24 (174–184) Author Location p24 (306–316 LAI) Epitope AEQASQDVKNW Subtype B Immunogen

**Species (MHC)** human (B\*4402, B44) **References** Brander & Walker 1997

• Pers. comm. from D. Lewinsohn to C. Brander and B. Walker, C Brander *et al.*, this database, 1999.

HXB2 Location p24 (174–184)
Author Location Gag (306–316)
Epitope AEQASQEVKNW
Immunogen HIV-1 infection
Species (MHC) human (B44)

**References** Brodie *et al.* 1999

- The ability of CTL effector cells was studied by expanding autologous HIV-1 Gag-specific CTL in vitro, and adoptively transferring them.
- The transferred CTLs migrated to the lymph nodes and transiently reduced circulating productively infected CD4+ T cells, showing that CTL move to appropriate target sites and mediate anti-viral effects.

HXB2 Location p24 (174–184)
Author Location p24 (306–316)
Epitope AEQASQEVKNW
Immunogen HIV-1 infection
Species (MHC) human (B44)
References Brodie et al. 2000

- Study tracks and quantifies in vivo migration of neo-marked CD8 HIV-specific CTL.
- Adoptively transferred gene-marked HIV-specific CTL homed to specific lymph node sites, colocalizing within the parafollicular regions of the lymph node adjacent to cells expressing HIV tat-fusion transcripts, indicative of viral replication.
- The CTL clones expressed CCR5 and localized among HIV-1 infected cells expressing MIP-1alpha and MIP-1beta, CCchemokines produced at sites of viral replication, suggesting a possible homing mechanism.
- This study provides a methodology for tracking and studying antigen specific CTL *in vivo*

HXB2 Location p24 (174–184)
Author Location p24 (306–316 LAI)
Epitope AEQASQDVKNW
Epitope name G3
Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B44)

Keywords HAART, ART

References Mollet et al. 2000

 A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.

- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened - eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+ PBL - but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

**HXB2 Location** p24 (174–184) Author Location p24 (174–184) Epitope AEQASQDVKNW Immunogen HIV-1 infection Species (MHC) human (B44) References Day et al. 2001

• B44-restricted CTL response was strongest to this epitope in one individual.

**HXB2 Location** p24 (174–184)

**Author Location** p24

Epitope AEQASQDVKNW **Epitope name** B44-AW11(p24)

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B44) Donor MHC A32, B44

> **Keywords** HAART, ART, supervised treatment interruptions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hivweb.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples of the 15 patients, and an additional 8 responses were detected only in LN. The total magnitude of the response was similar in LN and PB, but the percentage of CD8+ T cells in the LN is lower so the number of HIV-specific cells per million CD8+ T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- · Breakdowns of epitope responses were shown for 4 individuals. Patient B displayed the greatest response to epitope B44-AW11(p24) and also responded to A32-PW10(RT) in both PB and LN samples, while a third response against epitope A32-RW10(gp120) was only detected in the LN sample.

HXB2 Location p24 (174-184) Author Location Gag (B con) Epitope AEQASQEVKNW Epitope name AW11

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B44)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** variant cross-recognition crossneutralization

References Draenert et al. 2004c

- CD8+ T-cell responses that persisted in individuals with advanced disease were studied; most of the optimal epitopes defined were recognized with intermediate to high avidity. On average 13 (range, 2-39) epitopic regions were targeted in an average of 6 proteins (range, 1-8). HAART resulted in decrease in antigen and reduction in gamma IFN EliSpot responses, suggesting active responses to autologous virus. The lack of mutations within most viral epitopes suggest that persistent CTL through late disease do not exert strong immune selection pressure, yet the Elispot assays show robust responses, suggesting to the authors that gamma IFN-based screening methods may not reveal functional CD8+ T-cell impairment in patients with AIDS.
- 1 subject recognized this epitope with low functional avidity. The autologous sequence matched the B consensus.

HXB2 Location p24 (174-184)

**Author Location** (B consensus)

Epitope AEQASQDVKNW

Epitope name AW11

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B44)

**Donor MHC** A02, A11, B18, B44, Cw5, Cw12; A28, A29,

B14, B44, Cw8

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.
- 2/9 individuals recognized this epitope, presented by HLA-B44.

**HXB2 Location** p24 (174–184)

Author Location Gag (306–316)

Epitope AEQASQDVKNW

Epitope name AW11

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B44)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

**Keywords** optimal epitope **References** Allen *et al.* 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- This epitope did not vary.

**HXB2 Location** p24 (174–184)

Author Location Gag (306–316)

Epitope AEQASQDVKNW

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B44)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** p24 (174–184)

Author Location Gag (306-316)

Epitope AEQASADVKNW

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B44)

Donor MHC A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** p24 (174–184)

Author Location p24

Epitope AEQASQDVKNW

Subtype B, D

Immunogen HIV-1 infection

**Species (MHC)** human (B44) **Donor MHC** A23, A34, B44, B53, Cw4, Cw6

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype D Gag. The autologous epitope sequence had an D7E change, AEQASQeVKNW.

**HXB2 Location** p24 (174–185)

Author Location p24 (174–185)

Epitope AEQASQEVKNWM

Immunogen

Species (MHC) human (Cw5)

**Keywords** optimal epitope

References Frahm et al. 2007

**HXB2 Location** p24 (175–186)

**Author Location** p24 (307–318) **Epitope** EQASQEVKNWMT

Immunogen HIV-1 infection

Species (MHC) human (B44)

References Quayle et al. 1998

- HIV is found in semen both as cell-associated and cell-free forms, and HIV-specific CTL could be found in the semen of 5/5 men with CD4 greater than 500 3 of the men were analyzed in detail and had broad CTL to gag, env and pol.
- Two CTL lines from one donor recognized this epitope.
- Isolation of CTLs specific to HIV in both male and female urinal tracts provide evidence that virus-specific lymphocytes come from the urogenital mucosa, and the authors speculate that CTL in mucosal tissues may be correlated with lower viral load in semen and reduced transmission.

**HXB2 Location** p24 (176–184)

Author Location p24 (308–316 LAI)

Epitope QASQEVKNW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5301)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*5301 epitope.

**HXB2 Location** p24 (176–184)

**Author Location** (C consensus)

Epitope QATQDVKNW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- QATQDVKNW is an optimal epitope.

HXB2 Location p24 (176-184)

**Author Location** 

Epitope QASQEVKNW

Epitope name Gag-QW9

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5301, B57)

**Donor MHC** 01RCH59: A\*0201, A\*3201, B\*4002,

B\*5301, Cw\*0202, Cw\*0401

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- Subject 01RCH59 was Hispanic, was not on HAART, viral load 5100, CD4 count 349, and she also recognized PIQKETWETW, RT(392-401), A\*3201.
- Among HIV+ individuals who carried HLA B\*5301, 11/15 (73%) recognized this epitope.
- Among HIV+ individuals who carried HLA B57, 3/6 (60%) recognized this epitope.

**HXB2 Location** p24 (176–184)

Author Location p24 (309–317 LAI)

Epitope QASQEVKNW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

References Goulder et al. 1996b

- Recognition of this peptide by two long-term non-progressors.
- Peptide defined on the basis of B\*5801 binding motif, yet not cross-restricted except at high concentrations.
- Described as B\*5701 in C. Brander et al., this database, 1999.

**HXB2 Location** p24 (176–184)

**Author Location** p24 (311–319 LAI)

Epitope QASQEVKNW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

**Keywords** optimal epitope

**References** Frahm *et al.* 2007

• C. Brander notes this is a B\*5701 epitope.

**HXB2 Location** p24 (176–184)

**Author Location** 

Epitope QASQEVKNW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

**Keywords** rate of progression, immunodominance

References Migueles & Connors 2001

- HLA B\*5701 was found in a very high frequency in HIV-1 infected non-progressors, 11/13 (85%) versus 19/200 (9.5%) of progressors. Non-progressors tended to have an immune response that was highly focused on four p24 epitopes that were presented by B\*5701, ISPRTLNAW, KAFSPEVIPMF, TSTLQEQIGW, and QASQEVKNW.
- Only QASQEVKNW was recognized in all of the LTNP's tested.

HXB2 Location p24 (176–184)

**Author Location** 

Epitope QASQEVKNW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords rate of progression, immunodominance

References Migueles & Connors 2001

- CTL activity was monitored in 27 individuals, including 10 LTNP with an over-expression of HLA B\*5701 – these individuals have viral loads below the threshold of infection without therapy, and their CD8+ T-cell response tends to be focused on peptides that contain B\*5701 epitopes ISPRTLNAW, KAF-SPEVIPMF, TSTLQEQIGW, or QASQEVKNW.
- CTL responses are broader in B\*5701+ individuals with progressive viremia than those that control viremia.
- The HLA-A\*0201 SLYNTVATL epitope response was not as strong individuals that carried both A2, B57.

**HXB2 Location** p24 (176–184)

Author Location Gag (308–316)

Epitope QASQEVKNW

Epitope name QW9

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5701)

**Assay type** Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

Keywords rate of progression, escape

References Migueles et al. 2003

- cDNA Gag sequences from a set of 17 HLA-B\*5701+ progressors and 10 LTNPs were obtained, and the variation in four p24 B\*5701 epitopes examined. Sequence variants were more common (p < 0.01) in the epitopes in the progressors (median 3, range 1-7) than LTNPs (median 2, range 0-4).
- In general use of the autologous protein in a target cell did not diminish the overall CD8+ T-cell responses.
- The substituion E312D (qasqDvknw) was common in progressors (8/17) and rare in LTNP (1/8) (p = 0.06). qasqDvknw and qasqEvknw peptides were made; this mutation does not affect binding to B\*57. 2/4 progressors that carried only the D variant could not recognize the D variant peptide, but could recognize the E variant peptide, demonstrating immune escape.

 $\textbf{HXB2 Location} \hspace{0.1cm} p24 \hspace{0.1cm} (176\text{--}184)$ 

Author Location (C consensus)

**Epitope** QATQDVKNW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5801)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p24 (176–184)

Author Location Gag (308–316)

Epitope QASQEVKNW

**Epitope name OAS** 

Immunogen HIV-1 infection, HIV-2 infection

Species (MHC) human (B\*5801, B53)

Country Gambia.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords cross-presentation by different HLA, charac-

terizing CD8+ T cells

References Gillespie et al. 2005

- CD8 T-cell responses were evaluated and compared in HIV-1 and HIV-2 infected individuals. A significantly greater magnitude and breadth of Gag-specific responses were found in HIV-1 infected individuals, possibly because undetectable viral load in HIV-2 infected individuals. This study suggests that responses in HIV-2 infection reflect antigen load in plasma, as is the case in HIV-1 infection. No correlation was found between immune control of HIV-2 and the frequency of perforinexpressing virus-specific CD8 T-cells.
- QASQEVKNW is cross-presented by B53 and B\*5801. It was recognized in 4/4 B\*5801+ HIV-1 infected individuals, and 7/7 B53+ HIV-1 infected individuals. HIV-2 infected individuals preferentially recognized the B58 HIV-2 epitope TSTVEE-QIQW, and B53 epitope TPYDINQML.

HXB2 Location p24 (176-184)

Author Location p24 (308–316 LAI)

Epitope QASQEVKNW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B53)

References Buseyne et al. 1997

- Minimal sequence determined through epitope mapping.
- This is a relatively conserved epitope.

- HLA-Cw\*0401 was defined as the restricting element, but cells
  that carry Cw\*0401 varied in their ability to present this epitope
   this could be the result of diminished cell-surface expression
  of Cw\*0401 in some cells.
- The HLA presenting molecule for this epitope was originally described as Cw\*0401, but subsequent experiments with an HLA B53+ C4- cell line and with C1R cells transfected with HLA-B53 have shown that the HLA restricting element is HLA-B53 (F. Buseyne, pers. comm. 2000)

**HXB2 Location** p24 (176–184)

**Author Location** p24 (NL43)

Epitope QASQEVKNW

Epitope name QW9

Immunogen in vitro stimulation or selection

Species (MHC) human (B53)

Keywords epitope processing, dendritic cells

References Buseyne et al. 2001

- Exogenous presentation or cross-presentation of epitopes by antigen presenting cells (APC) without protein synthesis is an alternative pathway for CTL epitope processing that may be important in the initial generation of viral specific CTL.
- Dendritic cells treated with AZT to inhibit protein synthesis were able to elicit a strong specific CTL response in QASQEVKNW specific CTL clone 141 without protein synthesis, while macrophages demonstrated a decreased presentation efficiency.
- Exogenous Gag epitope presentation was Env-dependent and required receptor-dependent fusion.

**HXB2 Location** p24 (176–184)

**Author Location** p24 (308–316)

Epitope QATQEVKNW

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B53)

**Keywords** HIV exposed persistently seronegative (HEPS)

**References** Kaul *et al.* 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B53 women, 1/2 HEPS and 7/9 HIV-1 infected women recognized this epitope.

**HXB2 Location** p24 (176–184)

**Author Location** p24 (308–316 subtype A consensus)

Epitope QATQEVKNM

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (B53)

**Keywords** binding affinity, subtype comparisons **References** Dorrell *et al.* 2001

- In clade A infected Gambians, three HLA-B53 epitopes were defined in Gag p24 using ELISPOT, tetramer, and cytotoxicity assays.
- Two of the new epitopes lacked the predicted P2 anchors, DTI-NEEAAEW and QATQEVKNM, and bound to B53 with high affinity, thus extending the anchor residue motif for B53 and the related B35.
- While S, T, and P could all fit into the HLA-B35 or HLA-B53
  B pocket and form a hydrogen bond, A would not form a bond,
  so the authors propose compensatory interactions account for
  the high affinity of QATQEVKNM for B53.
- QATQEVKNM was recognized in 6/7 HLA-B53 subjects.
- Cross-recognition of QATQEVKNM was not studied here, but it was noted that both the A, QATQEVKNM, and B, QASQD-VKNW, subtype version of this epitope, are also presented by HLA-B57 and B58, common HLA alleles in Africans.

**HXB2 Location** p24 (176–184)

Author Location Gag (304-321 B con)

Epitope QASQEVKNW

Epitope name QW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B53, B58)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords variant cross-recognition or cross-

neutralization

References Draenert et al. 2004c

- CD8+ T-cell responses that persisted in individuals with advanced disease were studied; most of the optimal epitopes defined were recognized with intermediate to high avidity. On average 13 (range, 2-39) epitopic regions were targeted in an average of 6 proteins (range, 1-8). HAART resulted in decrease in antigen and reduction in gamma IFN EliSpot responses, suggesting active responses to autologous virus. The lack of mutations within most viral epitopes suggest that persistent CTL through late disease do not exert strong immune selection pressure, yet the Elispot assays show robust responses, suggesting to the authors that gamma IFN-based screening methods may not reveal functional CD8+ T-cell impairment in patients with AIDS.
- 2 subjects recognized this epitope with high functional avidity.
   Autologous sequence revealed no substitutions in this epitope compared to the B consensus.

**HXB2 Location** p24 (176–184)

**Author Location** Gag (SF2)

Epitope QASQEVKNW

Epitope name QW9

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords acute/early infection

References Goulder et al. 2001a

 This peptide elicited a weak CTL response during acute infection of patient PI004.  Three CTL responses, to epitopes TSTLQEQIGW, ISPRTL-NAW, and KAFSPEVIPMF, were evident early after infection; CTL responses to SLYNTVATL, QASQEVKNW, EIYKRWII, and FLKEKGGL were detectable at 5 months post-infection and beyond.

**HXB2 Location** p24 (176–184)

**Author Location** p24 (176–184)

Epitope QASQEVKNW

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 6/7 patients recognized this epitope.

HXB2 Location p24 (176–184)

**Author Location** Gag

Epitope QASQEVKNW

Epitope name QW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

Assav type CD8 T-cell Elispot - IFNγ

 $\label{lem:keywords} Keywords \ \ \text{subtype comparisons, escape, characterizing}$ 

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 5, qasqDvknw, was found in the most polymorphic residue in the epitope. This was shared between clades B and C. The most common substitution in people carrying B57 was in position 3, qaTqevknw.

**HXB2 Location** p24 (176–184)

**Author Location** p24 (308–316)

Epitope QATQDVKNW

Epitope name QW9

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Ethiopia.

Assay type CD8 T-cell Elispot - IFNγ

Keywords immunodominance, escape, variant cross-

recognition or cross-neutralization

References Currier et al. 2005

- Epitope sequence variation and CD8 T-cell responses were analyzed in C subtype infected HLA-B57-positive individuals from Ethiopia. KF11 was the immunodominant response.
- QATQDVKNW had a single variant, D5E (QATQEVKNW) in 1 subject; there was no apparent immune selection in this epitope. The QW9 peptide was tested in 2 B57-positive subjects; neither responded.

HXB2 Location p24 (176-184)

**Author Location** 

Epitope QASQEVKNW
Immunogen HIV-1 infection

Species (MHC) human (B57, B\*5801)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** responses in children, mother-to-infant transmission

References Feeney et al. 2005

• Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.

**HXB2 Location** p24 (176–184)

**Author Location** (LAI)

Epitope QASQEVKNW

Subtype B

Immunogen

**Species (MHC)** human (Cw4) **References** Buseyne 1999

**HXB2 Location** p24 (176–184) **Author Location** p24 (176–184)

Epitope QASGEVKNW

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (Cw4)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** p24 (176–184)

Author Location p24

Epitope QASQEVKNW

Subtype B, D

Immunogen HIV-1 infection Species (MHC) human (Cw4, B53)

Donor MHC A23, A24, B35, B58, Cw4, Cw7; A23, A34,

B44, B53, Cw4, Cw6

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from two infected people carrying subtype D Gag. The epitope sequence in one person matched the peptide, in the other had an E5D change, QASQdVKNW.

HXB2 Location p24 (176-184)

**Author Location** Gag

Epitope QASQEVKNW

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** assay standardization/improvement, epitope processing, characterizing CD8+ T cells

References Beattie et al. 2004

- This study compared CD8+ T cell EliSpot reponses to 58 Gag peptides that were optimal epitopes, with responses to overlapping 15 mers that spanned Gag. When screening for HIV-1-specific CD8 T-cell responses from 49 HIV+ people, overlapping 15-mer peptide pools revealed several novel responses that would have been missed using predefined CD8 epitopes. However, the 15-mer pools often missed low-level responses to predefined epitopes, especially when the epitope was located centrally in the 15-mer peptide, and the overall level of response to the 15 mers was generally lower(mean 1.4 fivel fold dilutions lower, range 0-3).
- In one individual, a response to QASQEVKNW could be detected at a concentration of 0.2 ug/ml, while a response to RAEQASQEVKNWMTE required 25 ug/ml for detection.

**HXB2 Location** p24 (176–185)

**Author Location** p24 (311–319 SF2)

Epitope QASKEVKNWV

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10

individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.

- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B57+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 0/0 group 2, and 1/2 group 3.

**HXB2 Location** p24 (177–185)

**Author Location** p24 (177–185)

**Epitope** ATQEVKNWM

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (B53)

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

- Variants A(T/S)QEVKNWM are specific for the A/B clades.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused
  on different epitopes with HLA presenting molecules that have
  previously been associated with reduced risk of infection, and
  there was a shift in the response in the HEPS women upon late
  seroconversion to epitopes recognized by the HIV-1 infected
  women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B53 women, 1/2 HEPS and 5/9 HIV-1 infected women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in the 1/2 HEPS case and in only one of the 5/9 HIV-1 infected women.

HXB2 Location p24 (180–189)

**Author Location** p24 (313–322)

**Epitope** EVKNWMTETL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B53)

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** p24 (181–190)

Author Location p24 (313–322 LAI)

**Epitope** VKNWMTETLL

Subtype B

Immunogen

Species (MHC) human (B8)

References Brander & Walker 1996

• P. Johnson, pers. comm.

HXB2 Location p24 (185-202)

**Author Location** (C consensus)

Epitope MTDTLLVQNANPDCKTIL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location p24 (191-205)

**Author Location** p24 (191–205)

Epitope VQNANPDCKTILKAL

Immunogen HIV-1 infection

Species (MHC) human (B51)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (191–205)

Author Location p24 (323–337)

Epitope VQNANPDCKTILKAL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Nixon & McMichael 1991

• Two CTL epitopes defined (see also p17(21-35))

**HXB2 Location** p24 (191–205)

**Author Location** p24 (325–339 SF2)

Epitope VQNANPDCKTILKAL

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords review, immunodominance, escape

References Goulder et al. 1997a; Phillips et al. 1991

- Longitudinal study of CTL escape mutants in people with the appropriate HLA types little variation was observed in the immunodominant B27 epitope, relative to the B8 epitopes, which varied over time.
- Goulder *et al.* [1997a] is a review of immune escape that points out that there may be a protective effect associated with B27, and that HLA-B8 individuals tend to progress more rapidly than HLA B27 patients.

**HXB2 Location** p24 (191–205)

**Author Location** Gag (320–328 BH10, LAI)

Epitope VQNANPDCKTILKAL

Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

 This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.  This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is TLLVQNANP) has similarity with growth differentiation factor 11, fragment THLVQQANP.

**HXB2 Location** p24 (191–210) **Author Location** p24 (323–342 SF2)

Epitope VQNANPDCKTILKALGPAAT

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- Three of these 12 had CTL response to this peptide.
- The responding subjects were HLA-A3, A24, B8, B55; HLA-A1, A11, B8, B27.

**HXB2 Location** p24 (191–210)

Author Location p24 (323–342 SF2)

Epitope VQNANPDCKTILKALGPAAT

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

 CTL expanded ex vivo were later infused into HIV-1 infected patients.

**HXB2 Location** p24 (193–201)

Author Location Gag (327–335 SF2)

Epitope NANPDCKTI

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

Keywords subtype comparisons, rate of progression

References Tomiyama et al. 1999

- HLA-B27, -B51, and -B57 are associated with slow progression to AIDS, while HLA-B35, -B8, -B24 are associated with a rapid progression to AIDS (Nat. Med. 2:405, 1996;Lancet 22:1187, 1986;Hum Immunol 22:73, 1988;Hum Immunol 44:156, 1995)
- 15% of Japanese populations carry HLA-B51 while HLA-B27 and -B57 are detected in less than 0.3%
- Of the 172 HIV-1 peptides with HLA-B\*5101 anchor residues, 33 bound to HLA-B\*5101, seven of these peptides were reactive with CTL from 3 B\*5101 positive individuals, and six were properly processed.
- Four of the six epitopes were highly conserved among B subtype sequences, NANPDCKTI is conserved.

**HXB2 Location** p24 (193–201)

**Author Location** p24 (193–201)

Epitope NANPDCKTI

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

**Donor MHC** A\*0201, A\*31, B\*27, B\*5101, Cw\*02; A\*2402, A\*26, B\*07, B\*5101, Cw\*07

Country Japan.

**Assay type** Chromium-release assay **Keywords** epitope processing, escape **References** Yokomaku *et al.* 2004

- Epitope variants escaped from being killed by CTLs in an endogenous expression system although they were recognized when corresponding synthetic peptides were exogenously loaded onto the cells. Escape is thus probably due to changes that occur during the processing and the presentation of epitopes in infected cells.
- Epitope variant nSnpdckNi was not recognized when added exogenously or when processed endogenously, but the mutations were in anchor residues and presumably inhibited binding to B\*5101.

HXB2 Location p24 (193-201)

Author Location p24 (325-333)

Epitope NANPDCKTI?

Immunogen HIV-1 infection

Species (MHC) human (B51)

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 3/11 of the HLA A2+ individuals were HLA B51 and two of these responded to this epitope as well as to other epitopes.

**HXB2 Location** p24 (193–201)

Author Location p24 (324–335 IIIB)

Epitope NANPDCKTI

Immunogen HIV-1 infection

Species (MHC) human (B51)

**Keywords** responses in children, mother-to-infant trans-

mission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- No variants of this epitope were found in a non-transmitting mother that had a CTL response to this epitope.

**HXB2 Location** p24 (193–201)

Author Location p24 (323-333)

Epitope NANPDCKTI

Epitope name NAN

Immunogen HIV-1 infection

Species (MHC) human (B51)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- None of the 8 study subjects recognized this epitope but none were HLA B51+

**HXB2 Location** p24 (193–201) Author Location p24 (193–201) Epitope NANPDCKTI Immunogen HIV-1 infection Species (MHC) human (B51) Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4, DR8, DQ7 Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- · Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

**HXB2 Location** p24 (193–201) Author Location p24 (193–201) Epitope NANPDSKTI Immunogen HIV-1 infection Species (MHC) human (B51) Donor MHC A\*3101, A68, B\*4403, B51 **Keywords** supervised treatment interruptions (STI)

References Arnedo-Valero et al. 2004

• T cell immune responses following STI were monitored in two chronically HIV-1 infected partners (A and B) who had contracted HIV-1 in 1992. STI induced strong transitory CD4+ and CD8+ T cell responses in both patients. Patient A displayed broad CD8+ T cell responses directed against Env, Pol, Gag, and Nef HIV-1 antigens. CTL responses in patient B were directed against two epitopes: Gag(p24)NANPDSKTI and Pol(RT)EELRQHLLRW.

**HXB2 Location** p24 (193–201)

Author Location p24

Epitope NANPDCKTI Immunogen HIV-1 infection Species (MHC) human (B51)

Donor MHC A01, A32, B\*1410, B15; A\*3101, A68, B\*4403, B51

Country Spain.

Assay type CD8 T-cell Elispot - IFNγ

Keywords HAART, ART, supervised treatment interruptions (STI)

References Arnedo-Valero et al. 2004

• T cell immune responses following STI were monitored in two chronically HIV-1 infected partners (A and B) who had contracted HIV-1 during 1992. STI induced strong transitory CD4+ and CD8+ T cell responses in both patients. The viruses remained very closely related over 10 years, despite the two individuals having different HLA types; the authors suggest the maintained similarity does not support a strong role for

HLA driven HIV diversity as has been claimed in Moore et al. (Science 2002).

• During the second treatment stop, patient A developed a strong proliferative response to p24, and multiple strong CD8+ T cell repsonses to Env, Pol, Gag and Nef. This patient was able to control viral load for two years follow up without therapy. Patient B developed a very weak CD4+ T cell response against p24 during breaks in therapy, and had CD8+ responses to two epitopes. Patient A: A01, A32, B\*1410, B15; Patient B: A\*3101, A68, B\*4403, B51.

**HXB2 Location** p24 (193–201) Author Location p24 (191–205) Epitope NANPDCKTI Immunogen HIV-1 infection Species (MHC) human (B8) References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** p24 (194–202) **Author Location** p24 (194–202) **Epitope** ANPDCKTIL **Epitope name** ANP Immunogen HIV-1 infection Species (MHC) human (B7) **Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401,

Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ

Keywords rate of progression, escape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant eptiopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This was one of 8 reactive epitopes found not to vary over time.

**HXB2 Location** p24 (195–202)

**Author Location** (C consensus)

**Epitope NPDCKTIL** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the T6 residue of NPDCKTIL are associated with the presence of the HLA presenting molecule in the host.

HXB2 Location p24 (195–202)
Author Location p24 (323–342)
Epitope NPDCKTIL
Immunogen HIV-1 infection
Species (MHC) human (B35)
References Bernard et al. 1998

- This study focuses on six rare long-term survivor HIV-infected people who were infected for many years without exhibiting immune dysregulation – such immunologically normal HIVinfected (INHI) cases occur at a frequency between 0.1 and 1% in the infected population.
- No direct CTL were found in any of the six INHIs, but above background CTLp activity was founded in 3/6 INHIs.
- Epitope sequences were deduced from larger reactive peptides based on HLA binding motifs – XPXXXXXL is a B35 binding motif.

**HXB2 Location** p24 (195–202) **Author Location** 

Epitope NPDCKTIL

Epitope name Gag-NL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA B35, 3/17 (18%) recognized this epitope.

HXB2 Location p24 (195–205) Author Location (C consensus) Epitope NPDCKTILRAL

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*3910) Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- NPDCKTILRAL is an optimal epitope.

HXB2 Location p24 (197–205)
Author Location p24 (329–337 LAI)
Epitope DCKTILKAL
Subtype B
Immunogen
Species (MHC) human (B\*0801)

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Keywords optimal epitope
References Frahm et al. 2007

• C. Brander notes this is a B*0801 epitope.
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c. Brander notes and is a B ooor epitope

HXB2 Location p24 (197–205) Author Location p24 (329–337 LAI) Epitope DCKTILKAL Subtype B Immunogen

Species (MHC) human (B8)

References Sutton et al. 1993

 Predicted epitope based on B8-binding motifs, from larger peptide VQNANPDCKTILKAL.

HXB2 Location p24 (197–205)
Author Location p24 (329–337)
Epitope DCKTILKAL
Immunogen HIV-1 infection
Species (MHC) human (B8)
Keywords escape

References Nowak et al. 1995

 In a longitudinal study of CTL response and immune escape – the variant DCRTILKAL was also found, binds to B8, but is not recognized.

HXB2 Location p24 (197–205)
Author Location p24 (329–337)
Epitope DCKTILKAL
Immunogen
Species (MHC) human (B8)

References McAdam et al. 1995

• Defined as minimal epitope by titration and binding studies.

HXB2 Location p24 (197–205) Author Location p24 (197–205) Epitope DCKTILKAL Immunogen

Species (MHC) human (B8)

References Goulder et al. 1997g

• Included in a study of the B8 binding motif.

HXB2 Location p24 (197–205)
Author Location p24 (329–337)
Epitope DCKTILKAL
Epitope name DCK
Immunogen HIV-1 infection
Species (MHC) human (B8)

**Keywords** HAART, ART, supervised treatment interruptions (STI), immunodominance, acute/early infection

References Oxenius et al. 2000

Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.

- This epitope was recognized at a low level by only 1 of the 7/8 study subjects that were HLA B8.

   HAART restores HIV specific immunity after advanced infection by increase of CD4+ and CD8+ T cell numbers after
- Patient SC12(HLA A1, B8/39, Cw0701/0702, DR2/3, DR51/52, DQ2/6) had sustained therapy started during acute infection and maintained an immunodominant response to FLKEKGGL throughout and minor responses to GEIYKR-WII, DCKTILKAL, GGKKKYKLK – GEIYKRWII and GGKKKYKLK responses were stimulated by a brief period off therapy.

Author Location p24 (197–205)

Author Location p24 (197–205)

Epitope DCKTILKAL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location p24 (197–205)
Author Location p24 (197–205)
Epitope DCKTILKAL
Immunogen HIV-1 infection
Species (MHC) human (B8)
References Day et al. 2001

• B8-restricted CTL accounted for about 1/3 of the total CTL response in one individual.

**HXB2 Location** p24 (197–205)

Author Location p24

Epitope DCKTILKAL

Epitope name DCK

Immunogen HIV-1 infection Species (MHC) human (B8)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius *et al.* [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there
  was no correlation between CTL responses with viral rebound
  rates, plateau viral loads, or clearance rates.

HXB2 Location p24 (197–205)
Author Location Gag (329–337)
Epitope DCKTILKAL
Immunogen HIV-1 infection
Species (MHC) humanized rabbit (B8)
Donor MHC A03, A28, B07, B08
Country Canada.

Assay type proliferation, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords HAART, ART, memory cells, immune dys-

function **References** Gamberg *et al.* 2004a

HAART restores HIV specific immunity after advanced infection by increase of CD4+ and CD8+ T cell numbers after supression of viral replication. However, HIV specific CTLs emerged only with detectable viral replication breakthroughs and were short-lived while CD4+ T-cell responses remained compromised, suggesting failure of generating stable CD8+ memory T-cells in the absence of HIV-specific T-helper responses.

HXB2 Location p24 (197–205)
Author Location (B consensus)
Epitope DCKTILKAL
Epitope name DL9
Subtype B
Immunogen HIV-1 infection

Species (MHC) human (B8)

Donor MHC A01, A03, B08, B14, Cw7, Cw8

Country United States.

Assay type Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** p24 (199–218) **Author Location** Gag (331–350)

**Epitope** KTILRALGPGATLEEMMTAC

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location p24 (203–211) Author Location Gag (335–343 SUMA)

Epitope KALGPAATL

Epitope name Gag KL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501, Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, acute/early infection, characterizing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4
   T cell counts through seven years of follow up. In contrast
   to more rapid progressors, WEAU and BORI, SUMA a broad
   response to 24 epitopes, with little immunodominance. Two
   peptides were somewhat more intensely recognized in acute
   infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

**HXB2 Location** p24 (209–217)

**Author Location** Gag (341–)

**Epitope** ATLEEMMTA

Epitope name Gag341

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: p24 Gag Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human, transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder that did not induce CTL and CD8+ T-cell IFN gamma responses in mice, although responses were detected in 2/17 HIV+ HLA-A2 subjects.

**HXB2 Location** p24 (211–230)

Author Location p24 (343–362 SF2)

Epitope LEEMMTACQGVGGPGHKARV

Immunogen HIV-1 infection Species (MHC) human (B7)

References McAdam et al. 1998

**HXB2 Location** p24 (211–230) **Author Location** p24 (345–364 SF2)

Epitope LEEMMTACQGVGGPGHKARV

Immunogen HIV-1 infection

Species (MHC) human

References van Baalen et al. 1993

 Gag CTL epitope precursor frequencies estimated, peptide mapping.

HXB2 Location p24 (211–231)

Author Location p24 (343–362 SF2)

Epitope LEEMMTACQGVGGPGHKARVL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI gag.
- One of these 12 had CTL response to this peptide.
- The responding subject was HLA-A1, A2, B50, B57.

HXB2 Location p24 (213-221)

**Author Location** Gag

**Epitope** EMMTACQGV

Epitope name E9V

Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV compo-

*nent:* gp140, gp140∆V3

**Species (MHC)** transgenic mouse (A\*0201)

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted
CTL responses against multiple conserved HIV-1 epitopes in
HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw
virus expressing gp140 with deleted V1, V2, and V3 loops
successfully induces neutralizing antibodies against HIV-1. A
live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination
against HIV and measles.

**HXB2 Location** p24 (213–221)

**Author Location** Gag (345–)

Epitope EMMTACQGV

Epitope name Gag345

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: p24 Gag Adjuvant: Incomplete Freund's Adju-

vant (IFA)

Species (MHC) human, transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay **Keywords** binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced a response in 1/6 transgenic mice. Responses were detected in 1/17 HIV+ HLA-A2 subjects.

**HXB2 Location** p24 (217–227) Author Location p24 (349–359 IIIB) Epitope ACQGVGGPGHK Immunogen HIV-1 infection Species (MHC) human (A\*1101) Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is an A\*1101 epitope.

**HXB2 Location** p24 (217–227) Author Location Gag (349–359) Epitope ACQGVGGPGHK Subtype B, CRF01\_AE Immunogen HIV-1 infection Species (MHC) human (A\*1101)

Keywords subtype comparisons, TCR usage

References Fukada et al. 2002

- Counterparts for eight known clade B HLA A\*1101 epitopes were generated for clade E (CRF01). Three epitopes, identical among clade A-E, were cross-reactive and recognized by clade E infected individuals. The clade E and B analogs to three more HLA A\*1101 epitopes was recognized in a clade specific manner. Two other HLA A\*1101 clade B defined epitopes were found not to have stimulated a response in clade E infected individuals.
- ACQGVGGPGHK was found to elicit clade-specific responses in clade B (ACQGVGGPGHK is most common in clades A and B) and clade E (acqgyggpShk is most common and is also common in clades C and D). ACQGVGGPGHK was recognized by CTL from 4/5 B clade infected Japanese subjects, and acqgvggpShk from 3/7 E clade infected Thai subjects.
- The binding of the two variants to HLA A\*1101 was almost identical, but bulk CTL generated from individuals did not cross-react with the cross-clade peptides, indicating the lack of cross-reactivity was due to TCR specificity.

**HXB2 Location** p24 (217–227)

Author Location Gag (349–359 SUMA)

Epitope ACQGVGGPGHK

Epitope name Gag AK11

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*1103)

B\*1501, Donor MHC A\*1103, A\*2402, B\*1402,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords dynamics, acute/early infection, characterizing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTLmediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4 T cell counts through seven years of follow up. In contrast to more rapid progressors, WEAU and BORI, SUMA a broad response to 24 epitopes, with little immunodominance. Two peptides were somewhat more intensely recognized in acute infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

HXB2 Location p24 (217-227) Author Location p24 (349–359 IIIB) Epitope ACQGVGGPGHK Immunogen HIV-1 infection Species (MHC) human (A11) References Sipsas et al. 1997

- · HIV IIIB proteins were used to define the range of CTL epitopes recognized by three lab workers accidentally infected with HIV-1 IIIB.
- · ACQGVGGPSHK, a variant found in HIV RF, was also recognized.

**HXB2 Location** p24 (217–227) Author Location p24 (SF2)

Epitope ACQGVGGPGHK

Immunogen HIV-1 infection

Species (MHC) human (A11)

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a • The CTL-dominant response was focused on this epitope in a HIV+ Caucasian living in Boston – this epitope did not fall

within the three most recognized peptides in the study.

- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPODLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

HXB2 Location p24 (217–227) **Author Location** p24 (349–359) Epitope ACQGVGGPGHK Epitope name ACQ

Immunogen HIV-1 infection Species (MHC) human (A11)

**Donor MHC** SC19: A\*11, A\*29, B\*08, B\*44, Cw\*06, Cw\*0701, DR3, DR7, DR52, DR53, DQ2, DQ8; SC18: A\*02, A\*11, B\*50, B\*58, Bw4, Bw6, Cw\*0401, Cw10, DR3, DR4, DR52,

DR53, DQ2, DQ8

**Keywords** HAART, ART, supervised treatment interruptions (STI), immunodominance, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- Both of the 2/8 HLA-A11 study subjects recognized this CTL epitope.
- Patient SC19(HLA A11/12, B8/44, Cw06/0701, DR3/7, DR52/53, DQ 2/8) had a CTL response to epitopes FLKEKGGL, GEIYKRWII, ACQGVGGPGHK, AVDLSH-FLK, and FNCGGEFFY that declined during therapy initiated at day 197.
- Patient SC18(HLA A2/11, B8/44, Cw06/0701, DR3/7, DR52/53, DQ2) recognizes the epitopes ACQGVGGPGHK, QVPLRPMTYK, AVDLSHFLK, and one called QIY but not fully described – he had brief therapy upon seroconversion and has had low viral load during 600 days of follow up.

Author Location p24 (217–227)

Author Location p24 (216–226)
Epitope ACQGVGGPGHK
Immunogen HIV-1 infection
Species (MHC) human (A11)
References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location p24 (217–227)

Author Location p24 (349–359 SF2)

Epitope ACQGVGGPGHK

Immunogen HIV-1 infection

Species (MHC) human (A11)

Keywords HAART, ART, acute/early infection

**References** Altfeld *et al.* 2001b Therapy provided during acute infection res

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.

- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A11+ individuals that had a CTL response to this epitope broken down by group: 0/3 group 1, 0/0 group 2, and 2/2 group 3.

**HXB2** Location p24 (217–227)

Author Location p24

Epitope ACQGVGGPGHK

Epitope name ACQ

Immunogen HIV-1 infection Species (MHC) human (A11)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius et al. [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there was no correlation between CTL responses with viral rebound rates, plateau viral loads, or clearance rates.

**HXB2 Location** p24 (217–227)

Author Location (B consensus)

Epitope ACQGVGGPGHK

Epitope name AK11

**Subtype** B **Immunogen** HIV-1 infection

Species (MHC) human (A11)

**Donor MHC** A11, A29, B08, B44, Cw4, Cw7

Country United States.

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** p24 (217–227)

Author Location Gag (349-359)

Epitope ACQGVGGPGHK

Epitope name AK11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, Flow cytometric T-cell cytokine

assay

or cross-neutralization, optimal epitope

References Allen et al. 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wild type, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wild type.
- A novel CD8 T-cell response was generated against the escape variant acqgvggpShk.
- · Additional analyses showed that the majority of individuals expressing HLA-A11 targeted the acggvggpShk variant sequence while the wild-type sequence was less frequently recognized.

**HXB2 Location** p24 (217–227)

**Author Location** Gag

Epitope ACQGVGGPGHK

Epitope name AK11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 9, acqgvggpShk, was found in the most polymorphic residues in the epitope. These were shared between clades B and C.

**HXB2 Location** p24 (221–231)

Author Location p24 (353–363 LAI)

Epitope VGGPGHKARVL

Epitope name G1

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords HAART, ART

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened – eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+ PBL - but with continued viral suppression, HIV-specific responses diminished.

**Keywords** escape. TCR usage, variant cross-recognition • Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

HXB2 Location p24 (223–231)

**Author Location (LAI)** 

Epitope GPGHKARVL

Subtype B

Immunogen

Species (MHC) (B\*0702)

Keywords optimal epitope

References Frahm et al. 2007; Goulder 1999

HXB2 Location p24 (223-231)

Author Location p24 (223–231 SF2)

Epitope GPGHKARVL

**Epitope name** GL9

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

References Altfeld et al. 2001a

- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- The response to GPGHKARVL was dominant.

**HXB2** Location p24 (223–231)

Author Location (C consensus)

Epitope GPSHKARVL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- · This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2** Location p24 (223–231)

**Author Location** (C consensus)

**Epitope** GPGHKARVL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the G3 residue of GPGHKARVL are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** p24 (223–231) **Author Location** (C consensus)

Epitope GPSHKARVL

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*0702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- GPSHKARVL is an optimal epitope.

**HXB2 Location** p24 (223–231)

**Author Location** (224–232)

Epitope GPSHKARVL

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*0702)

Assay type Other

Keywords HLA associated polymorphism

**References** Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- GPSHKARVL was a previously defined B\*0702 presented epitope that encompassed a B\*07- associated polymorphism, GPsHKARVL,in the third position.

**HXB2 Location** p24 (223–231)

Author Location p17

Epitope GPGHKARVL

Subtype D

Immunogen HIV-1 infection Species (MHC) human (B35)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7

**Country** Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

HXB2 Location p24 (223-231)

Author Location p24 (355–363 LAI)

Epitope GPGHKARVL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords review, escape

References Goulder et al. 1997e; Goulder et al. 1997a

- HLA-identical sibling hemophiliac brothers were both infected with the same batch of factor VIII.
- One had a strong response to this peptide, the other a weak response. They were tested 6-8 years after infection.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study.

**HXB2 Location** p24 (223–231)

**Author Location** p24 (SF2)

Epitope GPSHKARVL

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Keywords** subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in a HIV+ Caucasian living in Boston – this epitope did not fall within the three most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

HXB2 Location p24 (223-231)

Author Location p24 (SF2)

Epitope GPSHKARVL

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in a HIV+ Caucasian living in Boston – this epitope did not fall within the three most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

**HXB2 Location** p24 (223–231)

Author Location p24 (223–231 SF2)

**Epitope** GPGHKARVL **Immunogen** HIV-1 infection

Species (MHC) human (B7)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B7+ individuals that had a CTL response to this epitope broken down by group: 1/4 group 1, 2/3 group 2, and 0/1 group 3.

HXB2 Location p24 (223-231)

**Author Location** p24 (223–231)

Epitope GPGHKARVL

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Keywords** rate of progression, acute/early infection **References** Day *et al.* 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.

- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

**HXB2 Location** p24 (223–231)

**Author Location** p24 (223–231)

Epitope GPGHKARVL

**Epitope name** B7-GL9

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions

(STI), immunodominance, acute/early infec-

tion

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- Only two epitopes were detected during acute infection in patient AC-06, B7 restricted gp41 epitope IPRRIRQGL and Gag GPGHKARVL. GPGHKARVL was the first targeted peptide, and remained immunodominant through the 34 month study period.
- 3/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 1/4 individuals had detectable responses to this epitope after STI.

**HXB2 Location** p24 (223–231)

Author Location p24 (223–231)

Epitope GPGHKARVL

Epitope name B7-GL9 Gag

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

- An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the intial CTL response.
- In the earliest sample at day18 the sequence for this epitope was gpShkarvl. gpghkarvl dominated at day 606; both were equally well recognized.
- This was an immunodominant epitope, and was present in both viruses, the original strain and the superinfecting strain.

HXB2 Location p24 (223-231)

Author Location p24 (223–231)

Epitope GPGHKARVL

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFNγ, Flow

cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

• Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.

• 4/7 patients recognized this epitope.

HXB2 Location p24 (223-231)

Author Location (B consensus)

Epitope GPGHKARVL

Epitope name GL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A03, B07, Cw7

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

• Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.

• 1/9 individuals recognized this epitope.

**HXB2 Location** p24 (223–231)

Author Location Gag (355-363)

Epitope GPGHKARVL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

• Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades. • This epitope was reactive, but escape mutations did not accrue in it over time.

#### Gag p24-p2p7p1p6 CTL/CD8+ II-B-4 epitopes

HXB2 Location p24-p2p7p1p6 (223-1)

**Author Location** Gag

Epitope GPGHKARVLA

**Immunogen** 

Species (MHC) human (B7)

References De Groot et al. 2001

- The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.
- · A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes were identified that could stimulate IFNγ production in an ELISPOT assay.
- GPGHKARVLA was confirmed as an HLA-B7 epitope in this study, and had been previously published.

HXB2 Location p24-p2p7p1p6 (223-1)

**Author Location** Gag

Epitope GPGHKARVLA

Epitope name 1291

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A03, A24, B07, B38, Cw07, Cw12/13; A01,

A03, B07, B08, Cw03, Cw07

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for GPGHKARVLA: 28%

HXB2 Location p24-p2p7p1p6 (225-8)

Author Location Gag (357-372 LAI)

**Epitope** GHKARVLAEATLSQVN

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Buseyne et al. 1993a

- Vertical transmission of HIV ranges from 13% to 39%.
- Primary assays showed that cytotoxic activity against at least one HIV protein was detected in 70% of infected children.
- Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.

• Patient EM28 (CDC P2A) had a CTL response to four epitopes Author Location in Gag.

**HXB2 Location** p24-p2p7p1p6 (230–7)

Author Location Gag (386-)

Epitope VLAEAMSQV

Epitope name Gag-386

Immunogen

Species (MHC) human (A\*0201)

Keywords binding affinity, subtype comparisons, supertype, computational epitope prediction, immunodominance

References Altfeld et al. 2001c

- HIV was scanned for all peptides which carried the A2supermotif pattern conserved in more than 50% of B clade sequences - 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 - 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.
- Three additional previously described HLA-A2 epitopes were added to the set of 20, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- VLAEAMSQV binds to all five HLA-A2 supertype alleles tested: A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802 (highest affinity)
- 4/22 individuals with chronic HIV-1 infection recognized this epitope, and it was immunodominant in 3/4 by ELISPOT.
- 0/12 acutely infected individuals recognized this epitope.

HXB2 Location p24-p2p7p1p6 (230-7)

**Author Location** Gag

Epitope VLAEAMSQV

**Epitope name** Gag 386

Subtype M

Immunogen vaccine, in vitro stimulation or selection, com-

puter prediction

Vector/Type: DNA, peptide Adjuvant: In-

complete Freund's Adjuvant (IFA)

**Species (MHC)** human, mouse, humanized mouse (A\*0201)

Assay type Cytokine production, T-cell Elispot

**Keywords** subtype comparisons, computational epitope prediction, escape, TCR usage, variant cross-

recognition or cross-neutralization

References McKinney et al. 2004

- This study examined variant recognition of epitopes presented by A\*0201 and A\*1101. Numerous amino acid substitutions can be introduced into epitopes without disrupting their recognition by CTLs, although epitopes with multiple substitutions were less recognized. An algorithm was constructed for prediction of epitopes capable of inducing responses to a great number of variant epitopes.
- A total of 20 variant forms of Gag 386 were identified. More than 95% of the variant epitopes were recognized by CTLs from mice immunized with the parental form.
- Gag 386 epitope (parent or variant form) was present in 97% of HIV sequences of many M group subtypes.

HXB2 Location p24-p2p7p1p6 (230-7)

Epitope VLAEAMSQV

Epitope name Gag-VV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA A02, 3/29 (10%) recognized this epitope.

**HXB2 Location** p24-p2p7p1p6 (230–7)

**Author Location** Gag (362–)

**Epitope** VLAEAMSQV

Epitope name Gag362(9L)

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide Adjuvant: Incomplete

Freund's Adjuvant (IFA)

**Species (MHC)** human, transgenic mouse (A2)

Assay type T-cell Elispot, Chromium-release assay, Flow

cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder that induced CTL and CD8+ T-cell IFN gamma responses in mice. Responses were detected in 1/17 HIV+ HLA-A2 subjects.
- The variant vlaeamsqA was also immunogenic in A2 transgenic mice, eliciting a CD8+ T-cell response, as was recognized in 3/17 HIV+ people, including the person that recognized the vlaeamsqV variant.

**HXB2 Location** p24-p2p7p1p6 (230–7)

Author Location p24 (230–238)

**Epitope** VLAEAMSQV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

• The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized during chronic infection.

**HXB2 Location** p24-p2p7p1p6 (230–7)

**Author Location** p24-p2p7p1p6 (362–370)

**Epitope** VLAEAMSQV

Epitope name VV9

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A\*02, A\*32, B\*07, B\*40, Cw\*03, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection, reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs.
   Escape variants were commonly detected in maternal plasma.
   Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- The mother was A02+ and carried a variant form of the epitope, VLAEAMShV, which she passed to her A02- child. This form persisted in her child for 15 months.

**HXB2 Location** p24-p2p7p1p6 (230–7)

**Author Location** Gag (397–405)

**Epitope** VLAEAMSQV

Immunogen HIV-1 infection

Species (MHC) human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind all five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802)

### II-B-5 Gag p2p7p1p6 CTL/CD8 + epitopes

**HXB2 Location** p2p7p1p6 (1–10)

Author Location p2p7p1p6 (1-10)

**Epitope** AEAMSQVTNS

Immunogen

Species (MHC) human (B\*4501)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** p2p7p1p6 (5–13)

**Author Location** Gag (SF2)

Epitope SQVTNPANI

Immunogen vaccine

Strain: B clade SF2 HIV component: Gag

**Species (MHC)** mouse (H-2D<sup>b</sup>)

References Paliard et al. 1998

- HIV-1 (SF2)p55gag vaccination of H-2 mice activates a CTL response against this epitope.
- CTL that recognized SQVTNPANI in the context of H-2D<sup>b</sup> cross-reacted with H-2 alloantigens H-2L<sup>d</sup> and an unidentified self-peptide.
- A postulate: heterozygosity at the MHC level could prevent the maturation of some T cell receptor combinations for foreign peptide and self-MHC constructs because of thymic depletion and tolerance.

**HXB2 Location** p2p7p1p6 (8–17)

**Author Location** Gag

Epitope TNSANIMMQR

Epitope name TR10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 5, tnsaSimmqr, was found in the most polymorphic residues in the epitope. One escape mutation, at position 2,tTsanimmqr, was found not to correspond to the most polymorphic residues in the epitope. This is a novel partially mapped epitope.

**HXB2 Location** p2p7p1p6 (18–37)

Author Location Gag (96ZM651.8)

Epitope SNFKGNKRMVKCFNCGKEGH

Immunogen

Species (MHC) human (A\*020101)

References Novitsky et al. 2001

- This study is provides a survey of CTL responses and full length HIV-1 genome sequences from a C subtype infected Botswanan cohort.
- 4 of 8 individuals (50%) who were positive for HLA-A\*02011 responded to the peptide SNFKGNKRMVKCFNCGKEGH.

**HXB2 Location** p2p7p1p6 (24–31)

**Author Location** p6 (35–42 HXB2)

Epitope NSPTRREL

Epitope name NL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0102)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- Position4 in the epitope had potentially experienced positive selection. NsptRREL and tSPTRREL escape variants were found

HXB2 Location p2p7p1p6 (38-47)

**Author Location** Gag

Epitope LARNCRAPRK

**Epitope name** 1331 **Subtype** multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A03, A23, B49, B57; A03, A24, B27, B57, Cw13, Cw18; A03, A26, B08, B52

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for LARNCRAPRK: 35%. Immunodominant epitope.

**HXB2 Location** p2p7p1p6 (42–50)

Author Location p15 (42-50 SF2)

Epitope CRAPRKKGC

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

Donor MHC B14

Keywords immunodominance

References Yu et al. 2002b

- 26/57 HIV-1 infected subjects displayed Gag p15-specific CD8+ T-cell IFNgamma responses were measured by Elispot and intracellular staining. The immunodominant regions targeted by CD8+ T-cells were mapped to three functional domains: the zinc finger structures, the protease cleavage site p7/p1, and to the Vpr binding site in p6.
- p15 contributed on average 17% of the total Gag response (rage 0-100%).
- 3 optimal CTL epitopes were mapped within p15: KELY-PLTSL, CRAPRKKGC, and FLGKIWPSYK.
- 2/6 HLA-B14+ subjects recognized this epitope. The binding motif for B14 is C-term Cys, positions 2 and 5 Arg.

HXB2 Location p2p7p1p6 (42-50)

**Author Location** p15 (42–50)

Epitope CRAPRKKGC

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** p2p7p1p6 (42–50)

Author Location (B consensus)

Epitope CRAPRKKGC

Epitope name CC9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

Donor MHC A28, A29, B14, B44, Cw8

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope

HXB2 Location p2p7p1p6 (42–50)

Author Location Gag (405-413)

Epitope CRAPRKKGC

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** p2p7p1p6 (55–70)

Author Location p15 (446-460 BRU)

Epitope KEGHQMKDCTERQANF

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Claverie et al. 1988

• One of 4 epitopes first predicted, then subsequently shown to stimulate an HLA-A2 restricted CTL line.

HXB2 Location p2p7p1p6 (55–70)

**Author Location** Gag (41–56)

Epitope KEGHQMKDCTERQANF

**Immunogen** HIV-1 infection

Species (MHC) human (A2)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 19 patients recognized this epitope.

**HXB2 Location** p2p7p1p6 (58–69)

**Author Location** p24

Epitope HQMKDCNERQAN

Subtype B, G

Immunogen HIV-1 infection

**Species (MHC)** human (A2)

Donor MHC A2, A36, B45, B58, Cw3, Cw6

Country Nigeria.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, computational epitope

prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- This previously described epitope was embedded in a B clade peptide that was recognized by T-cells from an infected person carrying subtype G Gag. The autologous epitope sequence had 2 changes, an E8T substitution and a two amino acid insertion, QG at position 10, that could impact the anchor: HQMKDCNtRqgQAN.

**HXB2 Location** p2p7p1p6 (63–71)

**Author Location** p15 (63–71)

Epitope CTERQANFL

Immunogen HIV-1 infection

Species (MHC) human (B61)

Donor MHC A\*0201, A11, B51, B61, Cw2, Cw\*14

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

**HXB2 Location** p2p7p1p6 (64–71)

**Author Location** 

Epitope TERQANFL

Epitope name Gag-TL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*4002)

**Donor MHC** A\*0201, A\*3201, B\*4002, B\*5301,

Cw\*0202, Cw\*0401

**Keywords** HAART, ART **References** Sabbaj *et al.* 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- This epitope was newly defined in this study.
- Patient 01RCH59 was Hispanic, not on HAART, and had a viral load of 5100 and CD4 count of 349 she also recognized AEWDRVHPV, p24(78-86), HLA-B\*4002 and KEKGGLEGL, Nef(92-100), HLA-B\*4002.
- Among HIV+ individuals who carried HLA B40, 3/5 (60%) recognized this epitope.

**HXB2 Location** p2p7p1p6 (64–71)

**Author Location** p15 (64–71)

**Epitope TERQANFL** 

Immunogen

Species (MHC) human (B\*4002)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** p2p7p1p6 (66–74)

Author Location (C consensus)

Epitope RQANFLGKI

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*13) Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- RQANFLGKI is an optimal epitope.

 $\textbf{HXB2 Location} \hspace{0.1cm} p2p7p1p6 \hspace{0.1cm} (66\text{--}74)$ 

**Author Location** 

Epitope RQANFLGKI

Epitope name RI9

Immunogen

Species (MHC) human (B13)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B13 epitope.

**HXB2 Location** p2p7p1p6 (66–80)

Author Location p15 (66-80)

Epitope RQANFLGKIWPSYKG

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- Responses to this peptide were detected in 16% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** p2p7p1p6 (70–77)

Author Location Gag (433–)

Epitope FLGKIWPS

**Epitope name** Gag433

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Gag Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human, transgenic mouse (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder that induced CTL and CD8+ T-cell IFN gamma responses in mice. Responses were detected in 7/17 HIV+ HLA-A2 subjects.

**HXB2 Location** p2p7p1p6 (70–79)

Author Location p15 (70–79 SF2)

Epitope FLGKIWPSYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords immunodominance

References Yu et al. 2002b

- 26/57 HIV-1 infected subjects displayed Gag p15-specific CD8+ T-cell IFNgamma responses were measured by Elispot and intracellular staining. The immunodominant regions targeted by CD8+ T-cells were mapped to three functional domains: the zinc finger structures, the protease cleavage site p7/p1, and to the Vpr binding site in p6.
- p15 contributed on average 17% of the total Gag response (rage 0-100%).
- 3 optimal CTL epitopes were mapped within p15: KELY-PLTSL, CRAPRKKGC, and FLGKIWPSYK.
- FLGKIWPSYK was embedded in a peptide recognized by 14/57 (25%) of subjects.
- 13/24 (54%) of HLA-A\*0201+ subjects recognized this peptide

**HXB2 Location** p2p7p1p6 (70–79)

Author Location p2p7p1p6 (1-10)

**Epitope** FLGKIWPSYK

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** p2p7p1p6 (70–79)

**Author Location** (C consensus)

Epitope FLGKIWPSHK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Country South Africa.

**Assay type** CD8 T-cell Elispot - IFNγ **Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** p2p7p1p6 (70–79)

**Author Location** (C consensus)

**Epitope** FLGKIWPSHK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- FLGKIWPSHK is an optimal epitope.

**HXB2 Location** p2p7p1p6 (70–79)

**Author Location** (C consensus)

**Epitope** FLGKIWPSHK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0205)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- · FLGKIWPSHK is an optimal epitope.

**HXB2** Location p2p7p1p6 (70–79)

**Author Location** Gag (1–10)

Epitope FLGKIWPSYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) goat (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

• HLA class I restricted CD8+ T-cell responses against HIV-1 • The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized during chronic infection, in 54% of 74 chronically infected A2+ individuals, but in no acute cases (0/14).

HXB2 Location p2p7p1p6 (83-97)

Author Location p15 (418–433 BRU)

**Epitope** GNFLQSRPEPTAPPF

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Claverie et al. 1988

• One of 4 epitopes first predicted, then subsequently shown to stimulate an HLA-A2 restricted CTL line.

**HXB2 Location** p2p7p1p6 (83–97)

**Author Location** Gag (69–83)

**Epitope** GNFLQSRPTAPPF

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 19 patients recognized this epitope.

**HXB2 Location** p2p7p1p6 (83–97)

Author Location Gag (453-462 BH10, LAI)

Epitope GNFLQSRPEPTAPPF

Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is PEPTAPPFLQ) has similarity with the T-cell surface glycoprotein CD5, fragment PEPTAPPRLQ.

HXB2 Location p2p7p1p6 (91–100)

Author Location p2p7p1p6

**Epitope** EPTAPPEESF

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (B35, B58)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assav type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, computational epitope

prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

HXB2 Location p2p7p1p6 (108–116)

Author Location p2p7p1p6

Epitope TPSQKQEPI

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (B35)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, computational epitope

prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

**HXB2 Location** p2p7p1p6 (108–116)

Author Location p15

Epitope TPSQKQEPI

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (B53)

**Donor MHC** A23, A34, B44, B53, Cw4, Cw6

Country Democratic Republic of the Congo.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses,

which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.

• Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

**HXB2 Location** p2p7p1p6 (111–127)

**Author Location** Gag

Epitope QKQGTIDKELYPLASLK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This is a novel unmapped epitope. Two changes over time in the individual that recognized this peptide: QKQGTkDKELYPLvSLK

**HXB2 Location** p2p7p1p6 (113–121)

Author Location p15

Epitope QEPIDKELY

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (B44)

**Donor MHC** A23, A34, B44, B53, Cw4, Cw6

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

**HXB2 Location** p2p7p1p6 (114–123)

Author Location (C consensus)

Epitope EPKDREPL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- EPKDREPL is an optimal epitope.

**HXB2 Location** p2p7p1p6 (114–123)

Author Location p2p7p1p6

Epitope EPIDKELYPL

Subtype D

Immunogen HIV-1 infection Species (MHC) human (B35)

Donor MHC A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

- · Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

HXB2 Location p2p7p1p6 (114-123)

Author Location p15

Epitope EPIDKELYPL

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (B53)

**Donor MHC** A23, A34, B44, B53, Cw4, Cw6

Country Democratic Republic of the Congo.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

• Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.

• Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

**HXB2 Location** p2p7p1p6 (118–126)

**Author Location** Gag (481–489)

**Epitope KELYPLTSL** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*40)

Donor MHC A\*03, A\*24, B\*35, B\*40

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords acute/early infection, variant recognition or cross-neutralization, superin-

fection

References Yang et al. 2005b

- · An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- A response to this epitope was detected before superinfection but diminished afterward. The epitope in the infecting and superinfecting strain had the sequence: kelyplAsl. The second infecting strain had a 4-amino acid insertion proximal to the epitope, RGIDkelyplAsl.

**HXB2 Location** p2p7p1p6 (118–126)

**Author Location** p2p7p1p6 (118–126)

Epitope KELYPLTSL

Immunogen

Species (MHC) human (B\*4001)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is a B\*4001 epitope.

**HXB2 Location** p2p7p1p6 (118–126)

Author Location Gag p6 (481–489)

Epitope KELYPLTSL

Epitope name KL9

Immunogen HIV-1 infection

Species (MHC) human (B40)

**Donor MHC** A\*02, A\*32, B\*07, B\*40, Cw\*03, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords responses in children, mother-to-infant transmission, escape, acute/early infection, rever-

sion, viral fitness

References Sanchez-Merino et al. 2005

• CD8 T-cell responses were examined in mother-infant pairs. Escape variants were commonly detected in maternal plasma. Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype

and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.

 A B40+ mother carried the KEmYPLaSL variant of the epitope and transmitted it to her B40- infant. The variant form continued to dominate the infant's sequences at months 3 and 15.

**HXB2 Location** p2p7p1p6 (118–126) **Author Location** p15 (118–126 SF2)

**Epitope** KELYPLTSL

Epitope name p15-24

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B60)

**Keywords** immunodominance, cross-presentation by different HLA

References Yu et al. 2002b

- 26/57 HIV-1 infected subjects displayed Gag p15-specific CD8+ T-cell IFNgamma responses were measured by Elispot and intracellular staining. The immunodominant regions targeted by CD8+ T cells were mapped to three functional domains: the zinc finger structures, the protease cleavage site p7/p1, and to the Vpr binding site in p6.
- p15 contributed on average 17% of the total Gag response (rage 0-100%).
- 3 optimal CTL epitopes were mapped within p15: KELY-PLTSL, CRAPRKKGC, and FLGKIWPSYK.
- Four patients who were HLA-B60+ recognized KELYPLTSL.
- The binding motif for B60 is C-term Leu and 2nd position Glu.
- Four patients who did not carry HLA-B60 also recognized the 15 amino acid long peptide carrying KELYPLTSL, suggesting other epitopes in this immediate region can be presented by other HLA class I molecules.

**HXB2 Location** p2p7p1p6 (118–137)

**Author Location** Gag

Epitope KEMYPLASLRSLFGNDPSSQ

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Novel unmapped epitope.
- An S->L change was observed over time: KEMY-PLASLRSLFGNDPISQ

**HXB2 Location** p2p7p1p6 (120–129)

Author Location p2p7p1p6

**Epitope** LYPLASLRSL

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (A24)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope

prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

HXB2 Location p2p7p1p6 (121–129)

Author Location p15

**Epitope YPLASLRSL** 

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (B53)

Donor MHC A23, A34, B44, B53, Cw4, Cw6

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

HXB2 Location p2p7p1p6 (121–129)

**Author Location** p2p7p1p6 (36–44)

Epitope YPLASLRSL

Epitope name B7-YL9 Gag

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

- An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.
- The second infecting strain had the variant ypPlasIrsl. The CTL response was zero at all timepoints for the first variant. Insertion of a proline at position 3 (first variant) resulted in prevention of initial presentation of this region to the immune system.

**HXB2 Location** p2p7p1p6 (121–130)

**Author Location** p2p7p1p6

**Epitope** YPLASLRSLF

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (B35)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7 **Country** Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

HXB2 Location p2p7p1p6 (121–130) Author Location Gag (484–493) Epitope YPLTSLRSLF Immunogen HIV-1 infection

regions containing new epitopes.

Species (MHC) human (B7)

**References** Jin *et al.* 2000b

- This B7 epitope is one of three subdominant CTL responses detected in a long-term non-progressor.
- A dominant B7 epitope was defined using conventional methods, and three additional sub-dominant HLA B7 epitopes were defined by first using a non-anchor based strategy, EpiMatrix, to identify 2078 possible epitopes in the autologous HIV-1, followed by B7 anchor residue prediction to narrow the set to 55 peptides for experimental testing.

HXB2 Location p2p7p1p6 (121–130)

**Author Location** Gag

**Epitope** YPLTSLRSLF

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

**Donor MHC** A\*0301, A\*2301, B\*0702, B\*1503

Country United States.

**Keywords** escape, acute/early infection

References Bernardin et al. 2005

- Full HIV genomes from nine individuals were analyzed for mutations prior to seroconversion and 7 to 28 days later. It was found that the influence of the host's HLA type was reflected within weeks of infection; a statistically significant number of early nonsynonymous mutations were observed within previously reported CTL epitopes.
- T to A mutation was observed in position 4, and R to K in position 7.

HXB2 Location p2p7p1p6 (123–130)

Author Location p2p7p1p6

Epitope LASLRSLF

Subtype D

Immunogen HIV-1 infection

Species (MHC) human (B58)

**Donor MHC** A23, A24, B35, B58, Cw4, Cw7

Country Democratic Republic of the Congo.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction

References Geels et al. 2005

- Cross-clade CTL responses in 10 non-B HIV-1 infected individuals were evaluated. T-cell reactivity was tested against peptide pools of clade B Gag, Pol, Nef, Rev and Tat. Nine individuals demonstrated cross-reactive T-cell responses to clade A, B, and C Gag pools; 6/7 patients that responded to Nef-B also reacted to clade A and C Nef pools. An inverse correlation was observed between the sequence divergence of the HLA class I restricted epitopes and the height of the T-cell responses, which in Gag could be explained through variations in the HLA-anchor residues. 42% of the responses were directed to regions containing new epitopes.
- Predicted epitope based on an HLA motif embedded in a reactive peptide from a person carrying a D clade Gag.

## II-B-6 Gag CTL/CD8 + epitopes

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

Immunogen computer prediction

**Species (MHC)** (A\*0201, B\*3501)

**Keywords** subtype comparisons, computational epitope prediction

References Schönbach et al. 2002

 Computational methods (artificial neural networks, hidden Markov models, binding matrices based on HLA association rates) were used to identify HLA-A\*0201 and HLA-B\*3501 HIV T-cell epitope candidates from 533 Gag, Env and Pol sequences of which 374 were derived from HIV-1, 97 were derived from HIV-2 and 62 from SIV. Comparisons to known epitopes and between clades were made.

**HXB2 Location** Gag **Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection

**Species (MHC)** human (A\*0201, Cw\*08)

References Shacklett et al. 2000

HIV-1 specific, MHC class I-restricted CTL killing was detected in duodenal and rectal gut associated lymphoid tissue (GALT) sites from three infected individuals – the distribution of class I restricted CTL was different in the peripheral blood samples and GALT samples.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection Species (MHC) human (A2)

Assay type Tetramer binding, Flow cytometric T-cell cy-

tokine assay

Keywords assay standardization/improvement

References Wu et al. 2005

 A flow cytometric assay for validation of HIV-1 gag- or polspecific- CD8/HLA-A2 T-cells was shown to be sensitive and specific, being able to detect HIV-1 CTL at the single T-cell level. An inverse correlation between HIV plasma viremia and gag- and pol-specific-CD8/HLA-A2 T-cells was observed.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*35)

**Keywords** rate of progression

References Jin et al. 2002

- Patients with HLA-B\*35 variants B\*3502, B\*3503, B\*3504, and B\*5301 tend to proceed to AIDS more quickly than those with B\*3501.
- Of 32 patients with HLA-B\*35 alleles CD8+ CTL responses were quantified using an intracellular cytokine staining assay – 75% had responses to Pol, 69% to Gag, 50% to Nef, and 41% to Env.
- The overall magnitude of CTL responses did not differ between those bearing B\*3501 and the others. A higher percentage of Gag responses was observed in those that had lower RNA levels that carried B\*3501, and there was a negative association with viral load and CTL activity. The data is consistent with higher levels of CTL responses contributing to protection in B\*3501 individuals, but not in B\*3502, B\*3503, B\*3504, and B\*5301 individuals.

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

Subtype A, B, C

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost, canarypox prime with gp160 boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Gag, gp120, gp41,

Nef, Pol

Species (MHC) human (B60)

**Keywords** subtype comparisons, vaccine-specific epitope characteristics

References Ferrari et al. 2001

- Different HIV strains were used for different regions: gp41 LAI, Gag LAI, gp120 MN, gp120 SF2
- HLA-B60 responses dominated the responses against an Gag vaccine in an individual (022G0Z) who was HLA A1, A11, B8, B60. The strongest response was against the MN peptide 107-136. Low level Gag responses were observed against B8 and A11 epitopes, no response was observed against A1 epitopes.
- Vaccinee 202T7 (HLA A2, B27, C25) made the strongest response to an epitope at positions 131-140 of Gag. The response was highly cross-reactive with D clade Gag expressed from vaccinia, less so with C, and only minimally cross-reactive with A and CRF01.

**HXB2** Location Gag

Author Location p24

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component:

p17/p24 Gag

**Species (MHC)** mouse  $(H-2^b, H-2^d, H-2^k)$ 

References Iroegbu et al. 2000

- The p24 sequence is more conserved than is p17 within patient, and nonsynonymous substitutions are spread evenly throughout its coding regions, not concentrated in CTL epitopes.
- Minor changes in p24 did not alter the immunogenicity in H-2b,d, or k mice, while changes in p17 (92% similarity) did alter immunogenicity.

**HXB2 Location** Gag

**Author Location** Gag (SF2)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA, vaccinia Strain: B clade

SF2 HIV component: Gag, Pol

Species (MHC) mouse (H-2<sup>bxd</sup>)

References Otten et al. 2000

- CB6F1 were primed with gag DNA by im injection and challenged with vaccinia expressing Gag/Pol (rVVgag-pol)
- Gag-specific CTL responses were detected by IFNgamma secretion in the spleen, independent of the route (intraperitoneal, intranasal or intrarectal) of rVV gag-pol challenge.
- The gag DNA vaccine induced CTL responses in 4/4 monkeys 2 weeks post immunization, but antibody responses were detected in only 1/4 monkeys after 3 immunizations.
- CTL cross-reactivity against Gag sequences 1-80, 254-323, and 421-496 was observed, suggesting multiple CTL epitope recognition.

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Qiu et al. 2000

- Mice were injected with plasmid DNA at 0, 2 and 4 weeks and lymphocyte proliferation was measured after 6 weeks with recombinant p24 protein.
- Secreted HIV-1 Gag expression vectors generated a stronger response than standard Gag or cytoplasmic Gag expression vectors.
- IFN-gamma levels were increased compared to an undetectable IL-4 response.
- CTL levels were also increased in secreted Gag expression vaccination studies.

**HXB2 Location** Gag

Author Location Gag (SF2)

**Epitope** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade SF2

HIV component: Gag, Protease

**Species (MHC)** macaque, mouse (H-2<sup>d</sup>)

References zur Megede et al. 2000

- · Sequence-modified Rev-independent gag and gag-protease gene constructs lead to increased expression levels and elevated CTL and antibody immunogenicity in BALB/c and CB6F1 mice.
- · A CTL response in mice could be detected after a single immunization with codon-optimized gag, using 2 ng of plasmid; wild type gag required 200 ng to detect a response.
- Recognition of 3 different Gag peptide pools was observed, indicating a polyclonal CTL response.
- Significant gag-specific CTL responses were detected in 4/4 rhesus monkeys, in contrast to 1/4 using wildtype gag.

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen vaccine

Vector/Type: coxsackievirus HIV compo-

nent: p24 Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Halim et al. 2000

- An avirulent recombinant coxsackievirus (CB4-P) construct was generated that can express p24 Gag sequences – CB4-P is attenuated even in immunodeficient mice and T help responses can be elicited from peptides embedded in a surface loop of the VP1 capsid.
- This paper describes the vaccine strategy and generation of constructs, and employs amino-terminal fusion of Gag sequences to the viral polyprotein with subsequent cleavage to elicit CTL responses via MHC class I presentation in BALB/c mice.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade HXB2, B clade NL43 HIV component: Gag, Pol

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Huang et al. 2001

• Different HIV strains were used for different regions: gag HXB2, pol NL43

- Mice were immunized with four humanized DNA constructs: GagPol, that would form a pseudoparticle carrying Gag and Pol, Gag, Pol or a GagPol fusion construct.
- The GagPol pseudoparticle, Gag and GagPol fusion construct all elicited strong anti-Gag CTL, but only the GagPol fusion construct elicited strong anti-Pol CTL.

**HXB2 Location** Gag

Author Location Gag (HXB)

**Epitope** 

Immunogen vaccine

Vector/Type: Listeria monocytogenes Strain: B clade HXB2 HIV component:

Species (MHC) mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

Keywords Th1

References Mata et al. 2001

- BALB/c and C57BL/6 mice were immunized with recombinant Listeria monocytogenes (Lm-Gag) expressing HIV-1 HXB2 Gag and mice were challenged with vaccinia expressing Gag.
- L. monocytogenes is a gram-positive bacteria that enters the macrophage on phocytosis and lives in the cytoplasm - secreted L. monocytogenes antigens are processed and presented by both class I and class II pathways.
- CD4+ Th1 T-cells mediated the Gag specific immunological protection in mice immunized with Lm-Gag and challenged with vaccinia-Gag.
- · Gag-specific CTL may enhance viral clearance via IFN-gamma secretion, but are not essential for immunity.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen vaccine

Vector/Type: Listeria monocytogenes Strain: B clade HXB2 HIV component:

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

Keywords review, Th1

References Mata & Paterson 2000

- BALB/c and C57BL/6 mice were immunized with recombinant Listeria monocytogenes (Lm-Gag) expressing HIV-1 HXB2 Gag and mice were challenged with vaccinia expressing Gag.
- L. monocytogenes is a gram-positive bacteria that enters the macrophage on phagocytosis and lives in the cytoplasm - secreted L. monocytogenes antigens are processed and presented by both class I and class II pathways.
- This article is a review of L. monocytogenes biology and its potential as a vaccine vector for HIV, comparing to other vector systems, and discussing CD4+ Th1 T-cells mediated Gag specific immunological protection in mice and the Gag CTL response.

**HXB2** Location Gag

**Author Location** Gag (IIIB)

**Epitope** 

Immunogen vaccine

Vector/Type: virus-like particle (VLP) HIV

component: Gag

Species (MHC) macaque

References Paliard et al. 2000

- CTLs primed by HIV-1 p55 gag virus-like particle (VLP) vaccination recognized epitopes in four different 20 amino acid peptides p17/4, p17/8, p24/13 and p14/9.
- Cytotoxic T cell response lasted greater than 8.5 months.

**HXB2 Location** Gag

Author Location Gag (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression, Th1

References Wasik et al. 2000

- HIV+ infants that progressed rapidly to AIDS had lower Th1 responses and decreased production of beta-chemokines and IL-2 relative to other HIV+ infants.
- HIV+ infants had no demonstrable CTL at birth, but Th1 responses accompanied by CTL responses developed in children with slowly progressive disease, and not in rapid progressors.
- CTLp frequencies were determined by limiting dilution using autologous B cells infected with vaccina/HIV constructs.

**HXB2** Location Gag

**Author Location** Gag (LAI)

Epitope Subtype B

Immunogen vaccine

Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag, gp41. Protease, V3

Species (MHC) human

References Salmon-Ceron et al. 1999

- The vaccine used was a rec canarypox with HIV-1 gp120 MN, tm/gag/protease LAI (vCP205), alone or with p24E-V3 MN synthetic peptide (CLTB-36))
- Twenty HIV negative subjects were vaccinated in phase I trial with combinations of vCP205 and CLTB-36.
- Immunization with vCP205 induced HIV-1-specific ABs to gp120, V3, and p24 antigens, and CTL immune responses against vCP205 were detected after the fourth immunization in 33% of the subjects against Env, Gag and Pol, but the CLTB-36 peptide did not produce AB or CTL immune responses against p24 or gp160.

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen vaccine

Vector/Type: virus-like particle (VLP) HIV component: p17 Gag, p24 Gag

Species (MHC) human

References Klein et al. 1997

- Immunization of HIV+ people with an HIV-1 p17/p24 Ty viruslike particle (p24-VLP) resulted in a marginal, short-lived increased proliferative response to p24 and p17 and a transient elevation in viral load.
- Two of four subjects that received 500 or 1000  $\mu$ g of p24-VLP had an increase in gag-specific CTL.

**HXB2 Location** Gag

Author Location p24 (SF2)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade SF2 HIV component: gp120, p24 Gag Adjuvant: MF59, PLG

Species (MHC) mouse, baboon

References O'Hagan et al. 2000

PLG (Polylactide co-glycolide polymer) microparticles administered in MF59 emulsion induced gp120 Ab responses and CTL immune responses against p24 gag.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Lubaki et al. 1999

- Three strategies were used to analyze CTL activity: area under the net HIV-specific lysis curve (ACU), linear regression (LR) of net specific lysis, and the standard method, lytic units (LU20)
- A correlation between low HIV plasma viral load and increased levels of HIV-specific Gag and Nef CTL activity was observed using ACU and LR, but not LU20.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Kalams et al. 1999a

- The presence of HIV-1 p24-specific proliferative responses was positively correlated with Gag-specific memory CTL and negatively correlated with viral load in untreated subjects.
- Gag proliferative responses were the most readily detected –
  Gag CTL responses were the only responses with a significant
  correlation with Gag stimulated help, although there was a
  positive trend with Nef, Env and RT.

HXB2 Location Gag

Author Location p55 (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Greenough et al. 1999

 7/128 HIV-1 infected hemophiliac were identified as long-term non-progressors (LTNPs) and were monitored for viral and host immune parameters over 15 years – LTNPs maintained a low viral load, high frequencies of CTL precursors directed against Gag antigen and low levels of HIV-specific effector CTL activity – effector cell activity suggests low level ongoing viral replication.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Trickett et al. 1998

- Twelve HIV-1 infected patients were re-infused with their own lymphocytes, cryopreserved from an earlier time point in the infection.
- Improvement in CD4+ and CD8+ T cells was seen in 7/12, and an increase in the CTL response to Gag was seen in one patient.

HXB2 Location Gag

Author Location Gag (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Betts *et al.* 1999

This study demonstrated an inverse correlation between HIV
Type I plasma viral load and CTL activity directed against
HIV-1 Pol, and stronger combined effects of Pol- and Envspecific CTL, in long-term survivors (LTS) of HIV-1 infection.

**HXB2 Location** Gag

**Author Location** Gag (LAI)

Epitope

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Legrand et al. 1997

- Seventeen recently infected patients were tested for CTL response to HIV proteins Env, Gag, Pol, Rev, Nef, Vif and Tat.
- An early response (within a month following PI) was noted in 87% of the subjects to Gag, 75% to Env, and 50% to Nef.
- Early responses to Pol, Rev, Vif and Tat were rare.

**HXB2** Location Gag

Author Location Gag (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Betts et al. 1997

- 6/8 individuals from Zambia infected with C clade virus had CTL that were able to make response to B clade HIV-1 IIIB vaccinia-expressed Gag, Pol and Env proteins.
- A vigorous cross-clade response was not limited to a particular protein, and the level of recognition of different proteins varied among the six patients.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References De Maria et al. 1997

- CD3+ cells that also carry a natural killer cell receptor (NKR+) can exhibit down regulation of T cell function.
- Anti-NKR IgM MAb masked this inhibitory function and increased HIV-1 specific CTL activity in phytohemagglutininactivated PBMC cultured in the presence of IL-2 from 3/5 patients, and in one other case anti-NKR MAb brought HIV-1 specific CTL activity to detectable levels.

**HXB2** Location Gag

Author Location Gag (LAI)

**Epitope** 

Subtype B

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Gag, gp120, gp41. Protease

Species (MHC) human

References Belshe et al. 1998

 The live canarypox vaccine ALVAC-HIV(vCP205) carrying MN gp120, LAI gp41, Gag and Protease, and boosted with SF-2 rpg120, was given to HIV-1 seronegative volunteers – HIV-specific Env or Gag CD8+ CTL were detected in 64% of the volunteers.

**HXB2** Location Gag

Author Location Gag (LAI)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Buseyne et al. 1998a

 This study showed a correlation between strong CTL memory and breadth of response in 7-12 month old infants, and remaining AIDS-free for the first year of life, higher absolute CD4 and CD8 cells, and lower viral load.

**HXB2** Location Gag

**Author Location** Gag (LAI)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons

References Buseyne et al. 1998b

 In infants with positive CTL responses, most responses showed cross-clade reactivity with somewhat diminished recognition of epitopes from different subtypes.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

References Goh et al. 1999

- 13/37 exposed uninfected individuals with repeated high-risk sexual exposure had HIV-1 specific CTL against Env, Gag, Pol, or a combination of proteins CTL activity was correlated with a CCR5 wildtype genotype.
- In this group, the highest CTLp frequencies were directed at Gag, but the most common response was to Env and four individuals had responses to multiple HIV-1 proteins.

**HXB2 Location** Gag

**Author Location** Gag (LAI)

**Epitope** 

Subtype B

Immunogen vaccine

*Vector/Type:* canarypox *HIV component:* Gag, gp120, gp41, Nef, Protease, RT

Species (MHC) human

References Evans et al. 1999

 A Canarypox vaccine expressing gp120, gp41, Gag, Protease, Nef and Pol CTL epitopes gave rise to CTL that could be detected in 61% of the volunteers – responses to Gag, Env, Nef and Pol were detected 3-6 months after the last vaccination.

**HXB2 Location** Gag **Author Location** p17

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** epitope processing **References** Kuiken *et al.* 1999

- A correlation between conserved regions of p17 or Nef and CTL epitope density was noted. The authors suggest that this may be due to a biological reason such as epitope processing, or may be an artifact of experimental strategy for epitope definition, such that conserved epitopes would tend to be identified because they are more likely to be cross-reactive with the test reagents.
- In contrast to p17 and Nef, p24 is a more conserved protein, and known epitopes are evenly distributed across p24.

**HXB2 Location** Gag

Author Location Gag (LAI)

Epitope
Subtype B
Immunogen vaccine

Vector/Type: DNA prime with vaccinia boost Strain: B clade LAI HIV component: Env,

Gag

Species (MHC) macaque Keywords Th1, Th2 References Kent *et al.* 1998

- Priming with an HIV-DNA vaccine and boosting with a vaccinia construct induced greater levels of HIV T cell immunity than either vaccine alone.
- The proliferative response to Env and Gag after the DNA vaccination had a mean SI of 1.5-4, but after boosting with rHIV-fowlpox virus, there was a 6-17 fold increase in the mean SI for HIV Gag and Env. The T help response happened despite a fall in antibody titers, suggesting that the Th response was primarily Th1, not Th2. The CTL response was also enhanced.

HXB2 Location Gag

Author Location Gag/Pol (LAI, MN)

**Epitope** 

Immunogen vaccine

Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag, gp120, gp41, Protease

Species (MHC) human

References Salmon-Ceron et al. 1999

 A live attenuated canarypox vector expressing MN gp120 and LAI gp41/gag/protease could induce CTL and a lymphoproliferative response in healthy, uninfected volunteers. **HXB2 Location** Gag

Author Location Gag/Pol (MN)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env,

Gag, Pol Adjuvant: CD80, CD86

Species (MHC) chimpanzee

References Kim et al. 1998

 The study explores the use of co-stimulatory molecules coexpressed with an HIV-1 immunogen in a DNA vaccine to enhance the immune response – co-expression of CD86, but not CD80, dramatically increased both HIV Env and Gag/Pol specific CTL and Th proliferative responses.

**HXB2 Location** Gag

Author Location Gag (BRU)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression

References Aladdin et al. 1999

• *In vitro* measurements of CTL-activity by Cr release assay in bulk culture showed no correlation between CTL-activity (gp120, Gag, Pol and Nef) and disease progression as measured by viral load, CD4 and time to death.

**HXB2 Location** Gag

Author Location p24 (C consensus)

**Epitope** 

**Immunogen** HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons, immunodominance

References Goulder et al. 2000a

- The CTL-dominant response was focused on this epitope in a HIV+ South African this epitope did not fall within the five most recognized peptides in the study.
- Three peptides GSEELRSLYNTVATL (p17 residues 71-85), SALSEGATPQDLNTMLNTVG (p24 41-60), and WEKIRL-RPGGKKKYKLK (p17 16-30) contained the dominant Gagspecific epitope in 31 out of 44 B-clade infected individuals from Boston who showed Gag-CTL responses.
- Five peptides RLRPGGKKHYMIKHLVW (p17 20-36), ELRSLYNTVATLYCV (p17Gag 74-88), SALSEGATPQDL-NTMLNTVG (p24 41-60), FRDYVDRFFKTLRAEQA (p24 161-177), and SILDIKQGKEPFRDY (p24 149-164) contained dominant Gag-specific epitopes in 32 out of 37 C-clade infected subjects from South Africa.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: ZF1 HIV component: complete genome

Species (MHC) macaque

References Akahata et al. 2000

 Rhesus macaques were vaccinated by i.m. injection with naked plasmid DNA carrying an HIV-1 complete genome vaccine, strain ZF1, with a mutated zinc finger in the nucleocapsid to prevent packaging.

- Env and Gag specific CTL but no antibody responses were induced in 2/4 vaccinated monkeys (MM145 and MM153)
- 2/4 monkeys (MM146 and MM143) produced antibodies against p24 and/or gp160, but no CTL response was detected.
- PBMC from all vaccinated monkeys produced IFN-gamma, in response to HIV-1 gp160, indicating a Th response – this response was 5 times higher in MM145, the animal with the strongest CTL response.
- 4 weeks post-challenge with SHIV NM-3rN plasma viral loads of both MM145 and MM153 (with a homologous Env) decreased to near or below the detection limit.
- 6-8 weeks post-challenge with SHIV NM-3rN plasma viral loads of both MM146 and MM143 decreased near or below the detection limit.

HXB2 Location Gag Author Location Gag Epitope

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression

References Salerno-Goncalves et al. 2000

- A general test of CD8 anti-viral activity was developed based on proviral load of coculture of autologous CD8+ cells with CD4+ cells after homogeneous superinfection with NSI virus.
- Significantly decreased the CD4+ T-cell proviral loads were found in 12 HIV+ slow progressors relative to 10 rapid progressors.
- Significant CD8+ mediated cytotoxicity directed against autologous cells infected with vaccinia carrying the HIV-1 gag gene was observed in slow progressors in contrast to rapid progressors, but no correlation was found between plasma viral load in 22/22 asymptomatic HIV infected individuals.

HXB2 Location Gag Author Location Gag Epitope

Immunogen HIV-1 infection

Species (MHC) human

References Young et al. 2001

- Addition of recombinant rec human IL12 (rhIL12) to cultures increased HIV-specific lysis of HIV-Gag, Pol and gp120 vaccinia expressed antigens (11/15 tested increased lysis by > 5%) if the culture was derived from HIV+ individuals who had CD4 cells/ul > 500.
- 2/10 individuals with <200 CD4 cells/ul, and 3/10 individuals with 200-500 CD4cells/ul, had an increase of >5% upon treatment of the culture with rhIL12, so a few individuals in late stage disease had CD8 cells that maintained responsiveness to rhIL12.

HXB2 Location Gag Author Location p24 Epitope

Immunogen HIV-1 infection

Species (MHC) mouse

References de Quiros et al. 2000

CB-17 SCID-Hu mice engrafted with peripheral blood mononuclear cells of four long-term nonprogressors (viral load < 50 copies/ml) displayed resistance to challenge with HIV-1 SF162, mediated by CD8+ T-cells and associated with proliferation in response to p24 – these patients did not have a higher level of HIV-1 specific immunity *in vitro*, so the mechanism is unknown.

**HXB2 Location** Gag

Author Location Gag (subtype A, B, D)

**Epitope** 

Subtype A, B, D

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Cao et al. 2000

- HIV-1 subtypes A and D dominate the Ugandan epidemic, and a vaccine trial using B clade antigen is underway – this study addresses relative levels of cross-reactive CTL responses in HIV infected Ugandans to A, D, and B clade recombinant vaccinia viruses expressing Gag, Env, RT or Nef from HIV-1 clades A, B, and D.
- Proteins corresponding to the subtype of the infecting strains tended to trigger higher levels of CTL response measured by percent specific lysis, but there was extensive inter-subtype cross-reactivity with B clade proteins and the co-circulating subtype.

**HXB2** Location Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References White et al. 2001

 HIV-specific CTL activity was detected in the female reproductive tract of only 1/3 HIV-infected women who underwent a hysterectomy, although CTL could be identified in the PBMC of all three women.

**HXB2 Location** Gag

Author Location Gag (HXB2)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Chun et al. 2001

 Suppression of viral replication in the resting CD4+ T-cell reservoir by autologous CD8+ T-cells via CD4+/CD8+ cell contacts was observed in long-term nonprogressors and patients undergoing antiretroviral treatment, but this activity appears to be independent of Gag-specific CTL activity.

**HXB2 Location** Gag

**Author Location** Gag (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Jin *et al.* 2000a

- The CTL precursor level (CTLp) was measured in long term non-progressors (LTNP) with low viral load using limiting dilution analysis and measuring CTL against Env Gag and Pol expressed in vaccinia in autologous targets.
- LTNPs have high memory CTL numbers and low viral load.

**HXB2 Location** Gag **Author Location** Gag **Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

Keywords review, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 2001

- This is a review that summarizes observations about HIVspecific CTL found in the HIV-1 exposed persistently seronegative (HEPS) population.
- The CTL responses assayed by ELISPOT and by CTL precursor frequencies by limiting dilution analysis indicate that CTL in HEPS individuals tend to be of a lower magnitude than in chronic HIV-1 infections - the responses in HEPS cases are below the level of detection by tetramer assays.
- CD8+ CTL responses tend to be detectable in HEPS subjects only if they are recently exposed, and the response diminishes if exposure is reduced – it is not clear if there is a stable memory population in HEPS cases.
- CD8+ CTL responses in the HEPS population are associated with HIV-1 specific CD4+ T cell responses, assayed by proliferation assays, IL-2 secretion, and ELISPOT, and the authors consider the possibility that HIV-1-specific T-help responses improve the "quality" of the CD8+ response in HEPS individuals relative to HIV-1 infected individuals, who tend to have a poor HIV-1-specific T-help response.
- HIV-1 specific CD8+ CTL responses in HIV-1 infected individuals show reduced levels of perforin, and the T cells may not mature properly, and although similar studies have not been conducted in HEPS individuals this is considered as a possible difference in the CTL immune response in HEPS and HIV-1 infected people.

**HXB2** Location Gag

**Author Location** 

**Epitope** Subtype B

Immunogen vaccine

Gag, Pol

Species (MHC) mouse

Keywords review, vaccine-specific epitope characteristics

References Nabel 2002

• Using DNA that had humanized codon usage, CTL responses to DNA vaccines containing either Gag, Pol, Gag-Pol fusion protein, or Gag-Pol pseudoparticles suggested that the greatest breadth and most potent response was to the Gag-Pol fusion protein. The Gag-Pol fusion lacks the Gag precursor protein required for viral assemble, so does not form releaseable particles; the author speculates that longer retention of the Gag-Pol protein with in the cell may enhance antigen presentation.

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS), responses in children, mother-to-

infant transmission

References De Maria et al. 1994; Kuhn et al. 2002

- 6/24 HIV uninfected infants (ages 15-50 months) born to HIV+ mothers had HIV-1 specific CTL responses to vacciniaexpressed Nef, Gag/Pol, Env.
- Reviewed in Kuhn et al. [2002].

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords HIV exposed persistently seronegative (HEPS), responses in children, mother-to-

infant transmission

References Aldhous et al. 1994; Kuhn et al. 2002

- Six of nine HIV vertically infected infants had HIV-1 specific CTL responses to vaccinia expressed Tat (4/6), Pol (6/6), Env (1/6), or Gag (1/6), but not all responses were detected at all time points.
- Two of eleven babies that were not infected though born to HIV+ mothers had detectable responses to Tat (1/2), Pol (2/2),
- Reviewed in Kuhn et al. [2002].

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, HIV exposed persistently

seronegative (HEPS), responses in children, mother-to-infant transmission, rate of progres-

References Kuhn et al. 2002; Wasik et al. 1999

- Vector/Type: DNA HIV component: Env, In HIV-infected infants HIV-specific, CTL responses were not detectable in icord blood or in PBMC collected shortly after birth and were absent or remained very low in 3 infants with a rapidly progressive disease. For those who progressed more slowly, the HIV-specific CTL activity varied.
  - The two infants with high levels of Env peptide-stimulated IL-2 responses had the highest CTLp frequencies.
  - Stronger responses were detected after initiation of the antiretroviral therapy.
  - Two babies that were not infected though born to HIV+ mothers had detectable though low HIV-specific CTLp responses to Env (1/2), Pol (2/2), Gag (1/2) cord blood and transiently in PBMC after birth.
  - Reviewed in Kuhn et al. [2002].

**HXB2 Location** Gag

**Author Location** 

Epitope

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS), responses in children, mother-to-infant transmission

**References** Kuhn *et al.* 2002: McFarland *et al.* 1994

- Only 9% of HIV+ infants had HIV-specific CTL against Env or Gag in unstimulated PBMC. After CD3 stimulation of PBMC, Gag and Env specific CTL were found in PBMC from 91% and 78% of HIV-infected children, respectively, with high precursor frequencies.
- Reviewed in Kuhn et al. [2002].

**HXB2 Location** Gag

**Author Location** 

Epitope Epitope

Lpitope

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed. p17 is much more variable than p24.

**HXB2 Location** Gag

Author Location p24 (HXB)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, vaccine-specific epitope

characteristics

References Lu et al. 2000a

 Bacillus anthrax lethal toxin (LFn)-HIV fusion proteins are candidate HIV vaccines that are safe in mice, and LFn-V3 region fusion proteins induce CD8 T cells in BALBc mice. LFn causes exogenous protein to be taken up and processed in a class I pathway. Expressed proteins from Gag p24 and Nef fragments cloned into the LFn expression plasmid stimulate gag-specific CD4 proliferation and CTL responses in HIVinfected donor PBMCs in vitro.

HXB2 Location Gag

**Author Location** (HXB2)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Edwards et al. 2002

96% (26/27) chronically infected HIV-1 infected patients elicited gamma-IFN CD8+ T-cell responses against Gag.

- Nef and/or Pol CTL responses were detected in 86% of the subjects.
- The magnitude and breadth of Gag and p24 T-cell responses correlated with absolute CD4 counts, and inversely correlated with viral load.
- Pol and Int CTL responses correlated positively with absolute CD4+ T-cell count.
- Nef and Env responses did not correlate with either CD4 counts or viral load.

**HXB2 Location** Gag

**Author Location** 

Epitope

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, dendritic cells

References Larsson et al. 2002b

 Autologous mature dendritic cells with rec vaccinia expressing Gag, Pol, Nef and Env could amplify CD8+ T-cell Elispot responses 4-38 fold in five HIV+ patients on successful HAART treatment, relative to autologous monocytes. Some weak responses could only be detected using mature dendritic cells as APCs, and this approach could be useful for detection of low frequency memory cells.

**HXB2 Location** Gag

**Author Location** (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** immunotherapy

References Trickett et al. 2002

 Conditions were optimized for ex-vivo expansion of CD8+ and CD4+ T-cells with the goal of functional T-cell production for autologous immunotherapy. 10,000-fold expansions were obtained in 14 days with optimized concentrations of IL-2, anti-CD3 and anti-CD28 coated microspheres, and decreasing amounts of serum over the first 8 days.

HXB2 Location Gag

**Author Location (IIIB)** 

**Epitope** 

Subtype B

Immunogen HIV-1 and HCV co-infection

Species (MHC) human

Keywords rate of progression

References Lauer et al. 2002

- HIV-1 and HCV immune responses were studied in 22 individuals who were co-infected with HIV-1 and hepatitis C virus (HCV). IFNgamma production was measured in an Elispot assay of CD8+ T-cells using targets expressing either Gag, RT, Env and Nef in a vaccinia construct, or one of seven HCV proteins.
- All 22 patients targeted at least one protein. 20/22 patients recognized RT, 17/22 patients recognized Gag, 13/22 subjects recognized Env and 11/22 patients recognized Nef. Robust CTL activity was independent of disease progression or viral load.

- Despite high HCV viral loads, very few HCV CD8+ T-cell Elispot responses were detected. In a control HCV infected person who did not have HIV-1, strong anti-HCV responses were mounted.
- HIV-specific CD4 proliferative responses were detected in 9/17 coinfected patients, but no HCV responses were detected.

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** responses in children **References** Luzuriaga *et al.* 1995

- 2/3 infants infected in utero had detectable HIV-1 Gag and Env specific CTL responses, one by 4 months, one by 11 months of age. Levels of the responses varied at different time point. Pol responses were not detected.
- 2/4 infants infected intrapartum had detectable responses, one note until 11 months, one not until 42 months.
- HIV-specific CTL were not detected in ten HIV- infants that were born to HIV+ mothers.

HXB2 Location Gag

**Author Location** 

Epitope

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade LAI, B clade MN HIV component: Env, Gag

Species (MHC) human

References Gupta et al. 2002

- Different HIV strains were used for different regions: Gag, LAI; gp120, MN; and gp41, LAI
- A safety and immunogeniticity study of a vaccine dosing schedule was studied in a trial conducted in high and low risk study subjects. There was a 76% cumulative probability of detecting a Gag or Env CTL response by day 728.

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, responses in children

References Scott et al. 2001

- CTL responses before and after initiation of ART were studied in 13 HIV-1 vertically infected infants <6 months of age, and 4 that were >6 months of age.
- Before ART 2/13 infants <6 months of age showed IFNgamma Elispot CD8+ T-cell responses, one to Nef and one to Env and Nef, and these responses became undetectable after successful therapy—3 infants were coinfected with CMV and all 3 had CMV-specific CD8+ T-cell responses.
- One older infant, at 23 months, had CTL responses against all for proteins tested, Gag, Pol, Nef and Env, and had the lowest plasma viremia of the study group. 3/4 infants older than 6 months of age responded to either Nef or Pol.

 Administration of ART over 48 weeks broadened the HIV-1specific CTL response in 2/4 of the older children that were incomplete responders.

**HXB2** Location Gag

Author Location (IIIB, MN)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords dendritic cells

References Larsson et al. 2002a

 Dendritic cells acquire and present HIV-1 antigens derived from dead, apoptotic cells or from non-infectious, fusioncompetent HIV-1 virions, and these DC cells could stimulate CD4+ and CD8+ T-cells resulting in IFNgamma production in an Elispot assay. Both HLA Class I and class II molecules were used for presentation. This may be an important aspect of the initial immune response to HIV-1 infection of CD4+ cells in the mucosal subepithelia.

**HXB2 Location** Gag

**Author Location (IIIB)** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Ortiz et al. 2001

 Immune responses in eight chronically HIV-1 infected patients undergoing HAART therapy structured treatment interruptions (STI) were studied. STI boosted HIV-1 specific CTL responses and elevated CTL responses were maintained up to 22 weeks after the last treatment interruption, but viral load rebound to pretreatment levels and CD4 T-cell count decline was observed. CD8 responses in PBMC were measured by cytokine flow cytometry with gp160, Gag p55, RT-Pol and Nef expressed in vaccinia.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Subtype AG, B

Immunogen

Species (MHC) human

References

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen

Species (MHC) human

References

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type Intracellular cytokine staining

**Keywords** HAART, ART, computational epitope prediction, supervised treatment interruptions (STI)

References Amicosante et al. 2002

- A new assay was developed to detect CTL responses to HIV using 28 pooled 15-mer peptides from conserved regions in Gag that were selected to be rich in HLA class I motifs, carrying potential epitopes for more than 90% of HLA class I haplotypes, and to be conserved between subtypes. Some peptide variants were included, expanding the potential for cross-clade recognition. 12 Caucasians, even those on successful HAART, had detectable CTL responses using this assay, and as did five Africans. People with either B subtype or A-G recombinant infections all reacted.
- The Gag peptide ICS assay was more sensitive to picking up CTL reactivity than whole Gag in HAART treated people. Initiation of STI increased the number of IFN-gamma producing CD8+ T-cells detected using the peptide assay.

HXB2 Location Gag
Author Location Gag
Epitope

Immunogen vaccine

Vector/Type: vaccinia HIV component: Gag Adjuvant: block copolymer CRL8623

Species (MHC) macaque

**Assay type** CD8 T-cell Elispot - IFN $\gamma$  **Keywords** vaccine-induced epitopes **References** Caulfield *et al.* 2002

- Codon-optimized HIV Gag DNA vaccines were given i.m. with or without a nonionic block copolymer(CRL8623) as adjuvant. DNA-CRL8623 fourmulations induced 2-fold higher Elispot responses, shifting the response towards CD8+ T-cells.
- 23 monkeys recognized 25 different epitopes with an average of 2.7 epitopes per monkey, and a minimum of 1 and a maximum of 5 peptides per monkey.
- Responses were detected up to 18 months after vaccination.

HXB2 Location Gag
Author Location Gag
Epitope
Subtype multiple

Immunogen
Species (MHC) human

Assav type Flow cytometric T-cell cytokine assay

**Keywords** subtype comparisons

References Currier et al. 2003

- CD8-cellular immune responses from 21 HIV-1 infected patients from Kenya infected with subtype A, C, D, and unique recombinants were studied for cross-recognition of Gag, Env, and Nef vaccinia-expressed proteins representing subtypes A-H, including CRF01.
- Both subtype-specific and cross-reactive CTL were observed, with a skewing of responses towards the infecting subtype in the nine subjects with full length sequence available. The magnitude of the responses to Gag were the highest, less to Nef, and still less to Env.
- For Gag, 8/21 subjects responded to at least 7/8 different subtype proteins, 7 had a mixed response, recognizing some subtypes but not others, and 6 responded to only one or none of the different subtype proteins. For Env, 4/19 subjects responded to

at least 7/8 subtypes, 8 had a mixed response, and 7 responded to one or none. For Nef, 5/19 subjects responded to at least 7/8 subtypes, 8 had a mixed response, and 6 responded to one or none.

HXB2 Location Gag Author Location Gag (SF2) Epitope

Subtype B

Immunogen vaccine

Vector/Type: DNA, protein, virus-like particle (VLP), PLG microparticle Strain: B clade SF2 HIV component: Gag Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72), LTK63

Species (MHC) macaque

**Assay type** proliferation, Chromium-release assay **References** Otten *et al.* 2003

Immunization strategies for Gag (p55) in macaques were compared. GAG DNA prime with a boost of Gag adsorbed onto PLG (polyactide coglycolide) microparticles with LTK63 as adjuvant gave the strongest CD4+ T cell proliferative, CTL, and antibody responses, compared with Gag protein, or Gag viruslike particles (VLP). GAG DNA was best for inducing CTL responses, Gag-PLG for T-help and antibody; the prime-boost combination gave strong responses for all three.

**HXB2 Location** Gag **Author Location** Gag

**Epitope** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, immunodominance, acute/early infection, early-expressed proteins

References Masemola et al. 2004a

- Anti-HIV T-cell responses in subtype C HIV-1 infected individuals in the beginning of the infection target multiple protein regions, but the responses are dominated by Nef, making up almost one-third of the total responses; the second most targeted protein was p24. A correlation between Gag specific responses and plasma viral load was also found.
- Neither breadth nor magnitude of CD8+ T-cell responses were correlated with control of virus, however hierarchical preferential targeting of Gag was significantly associated with lower viral loads.

**HXB2 Location** Gag

**Author Location** Gag

Epitope

Subtype A

Immunogen vaccine

Vector/Type: DNA, modified vaccinia Ankara (MVA), polyepitope, DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17/p24 Gag

Species (MHC) human

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFNγ

Keywords epitope processing, vaccine-induced epitopes,

vaccine antigen design

References Mwau et al. 2004

- Phase I clinical trial in healthy uninfected individuals was conducted evaluating the immunogenicities of candidate DNA- and MVA-vectored HIV vaccines. Both DNA and MVA vaccines alone and combined (DNA prime-MVA boost) were shown to be safe and induce HIV-specific responses in 78%, 88% and 89% of individuals, respectively. Responses in some individuals could be detected 1 year after vaccination.
- The vaccine in this case was a clade A p17/p24 antigen linked to a polyepitope string of A clade eptiopes. Reponses were tested with peptide pools, and multiple strong responses to the gag proteins and to the poylepitope region were observed. MVA alone did as well as a DNA prime, MVA boost in this study, although the study included small numbers.

**HXB2 Location** Gag

**Author Location** Gag

Epitope Subtype B

Immunogen vaccine

Vector/Type: non-replicating adenovirus

Strain: B clade HIV component: Gag

Species (MHC) mouse

Assay type Intracellular cytokine staining

**Keywords** Th1, Th2 **References** Pinto *et al.* 2003

 Heterologous prime boosts with replication-defective adenoviral vectors of different simian serotypes expressing the same transgene product of HIV-1 were shown to be highly efficient in increasing specific CD8+ T-cell responses.

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

Subtype CRF02\_AG

Immunogen vaccine

Vector/Type: virus-like particle (VLP), DNA prime with modified vaccinia Ankara (MVA) boost Strain: CRF02 IC0928 HIV compo-

nent: Env, Gag, Pol

Species (MHC) macaque

**Assay type** T-cell Elispot, Intracellular cytokine staining **Keywords** vaccine-specific epitope characteristics, vac-

cine antigen design

References Ellenberger et al. 2005

 Macaques were given a Gag-Pol-Env DNA prime followed by an MVA boost. Two DNA constructs were compared, one that resulted in mature VLPs with processed Gag (IC48) and one that had a point mutation in Gag that resulted in immature VLPs (IC1-90). IC48 DNA vaccinations, which produced mature VLPs, yielded 2-fold stronger T-cell responses with greater breadth. CD4 T-cells responded to 3-fold more peptide pools than did CD8.

HXB2 Location Gag Author Location Gag Epitope Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Country South Africa.

Assay type Flow cytometric T-cell cytokine assay

Keywords characterizing CD8+ T cells

References Ramduth et al. 2005

 The magnitude of HIV-specific CD8 responses in HIV-1 infected individuals from South Africa correlated with the CD4 responses. CD4 responses were much more narrowly focused, with Gag as dominant target, while CD8 responses were equally distributed among Gag, Pol and the regulatory and accessory proteins. An association between the preferential targeting of Gag by CD8 T-cells and viral control was found.

**HXB2** Location Gag

Author Location Gag (Consensus B, DU422)

**Epitope** 

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** assay standardization/improvement, subtype comparisons, variant cross-recognition or

cross-neutralization

References Sabado et al. 2005

 CD8 T-cell responses were tested in HIV-1 clade B infected individuals using Gag peptides based on clade B consensus sequence and clade C primary isolate DU422. Peptides from both clades were shown to be of equal sensitivity, with equal numbers of discordantly detected responses. The majority of discordant detection was due to sequence differences between clades. Thus, clade B consensus peptides were not superior in detecting CD8 T-cell responses in clade B-infected individuals.

# II-B-7 Gag/Pol CTL/CD8 + epitopes

HXB2 Location Gag/Pol

Author Location Gag/Pol (ARV-2 SF2)

Epitope

Immunogen vaccine

Vector/Type: fowlpoxvirus Strain: B clade ARV-2, B clade SF2 HIV component: Gag,

Pol Adjuvant: IFNγ

Species (MHC) macaque

References Kent et al. 2000

- Vaccination with FPV Gag/Pol-IFN-gamma increased HIV-1 specific CTL and T cell proliferative responses to Gag/Pol antigens, respectively, in infected Macaca nemestrina.
- HIV-1 viral loads remained low and unchanged following vaccinations.

HXB2 Location Gag/Pol Author Location RT Enitone

Epitope Immunogen vaccine Vector/Type: DNA HIV component: Env, II-B-9
Gag. Pol. Vif Adjuvant: B7, IL-12

Species (MHC) mouse

References Kim et al. 1997d

- A Gag/Pol or Env DNA vaccine, when delivered in conjunction with the plasmid encoding the co-stimulatory molecules CD86, gave a dramatic increase in both the cytotoxic and proliferative responses in mice.
- When CD86 was present, CTL response could be detected even without *in vitro* stimulation.

HXB2 Location Gag/Pol

Author Location RT

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords TCR usage

References Gamberg et al. 1999

- 13/13 subjects with advanced HIV infections showed CD8 T cell proliferation and differentiation of CTL *in vitro*, and six individuals showed HIV-specific responses to Gag, Pol, Env or Nef antigens.
- Data suggests that the functional and genetic integrity of the CD8 T cell repertoire (TCR Vβ gene intrafamily genetic diversity) remains intact through advanced HIV infection, although HIV-specific CTL activity decreases.

HXB2 Location Gag/Pol

**Author Location** 

**Epitope** 

Immunogen vaccine

Vector/Type: adenovirus HIV component:

Gag-Pol, Nef, Vpr

Species (MHC) mouse

References Muthumani et al. 2002

- Vpr can cause cells to go into G2 arrest, and it surpresses immune cell activation and inflammatory cytokine production, so co-immunization of BALB/c mice with recombinant adenovirus expressing Vpr and HIV-1 antigens Nef or Gag/Pol was tested to see if Vpr reduced the immune response to the other HIV antigens.
- Vpr compromised CD8+ T-cell lytic responses and T-helper proliferative responses in mice co-immunized with Vpr and Nef or Gag/Pol.
- In vitro, Vpr reduced T-cell cytokine production of IL-12 and TNFalpha, indicative of Vpr-mediated immune suppression.

## II-B-8 Gag/Pol TF CTL/CD8+ epitopes

HXB2 Location Gag/Pol TF (24-31)

**Author Location** 

Epitope NSPTRREL

Epitope name NL8

Immunogen

Species (MHC) human (Cw\*0102)

**Keywords** optimal epitope

**References** Frahm *et al.* 2007

• C. Brander notes this is a Cw\*0102 epitope.

# II-B-9 Gag/Pol TF-Protease CTL/CD8 + epitopes

HXB2 Location Gag/Pol TF-Protease (54-6)

**Author Location** Pol

Epitope FSFPQITLW

**Epitope name** FW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- An escape mutation at position 2, fNfpqitlw, was found in the most polymorphic residue in the epitope. This is a novel partially mapped epitope.

## II-B-10 Protease CTL/CD8+ epitopes

HXB2 Location Protease (2–19)

**Author Location** (C consensus)

Epitope QITLWQRPLVSIKVGGQI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0801)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Protease (3–11)

Author Location RT (71-79 subtype A, B, D)

Epitope ITLWQRPLV

Subtype A, B, D

Immunogen

Species (MHC) human (A\*6802)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*6802 epitope.

**HXB2 Location** Protease (3–11)

**Author Location** Pol

Epitope ITLWQRPLV

Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade

HIV component: p17 Gag, p24 Gag

Species (MHC) human (A\*6802)

Keywords subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location Protease (3-11)

**Author Location** Protease (71–79 LAI)

**Epitope ITLWQRPLV** 

Subtype B

Immunogen

**Species (MHC)** human (A\*6802, A\*7401, A19)

**Keywords** subtype comparisons

**References** Dong 1998

- Predicted on binding motif, no truncations analyzed.
- Clade A/B/D consensus, S. Rowland-Jones, pers. comm.

**HXB2 Location** Protease (3–11)

**Author Location** RT (71–79 subtype A, B, D)

**Epitope ITLWQRPLV** 

Subtype A, B, D

Immunogen

Species (MHC) human (A\*7401)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*7401 epitope.

**HXB2 Location** Protease (3–11)

**Author Location** Pol (59–)

Epitope ITLWQRPLV

Epitope name Pol59

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Protease Adjuvant: Incomplete Freund's Adju-

vant (IFA)

Species (MHC) transgenic mouse (A2)

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was an intermediate A2 binder, and induced CTL and CD8+ T-cell IFN gamma responses in mice. Responses to the peptide were not detected in 17 HIV+ HLA-A2 subjects.

HXB2 Location Protease (3-11)

**Author Location** Pol (59–65)

Epitope ITLWQRPLV

Immunogen HIV-1 infection

Species (MHC) human (A28)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** Protease (3–11)

**Author Location** Pol (60–68)

**Epitope ITLWQRPLV** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A28)

Donor MHC A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

HXB2 Location Protease (3-11)

Author Location RT (71-79 LAI)

Epitope ITLWQRPLV

Epitope name P2

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (A28 supertype)

**Keywords** HAART, ART, supertype

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened - eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+

PBL – but with continued viral suppression, HIV-specific responses diminished.

 Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

HXB2 Location Protease (3-11)

Author Location Pol

Epitope ITLWQRPLV

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A74)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

• ITLWQRPLV cross-reacts with clades A, B and D.

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** Protease (4–14)

**Author Location** Pol (60–70 SF2)

Epitope TLWQRPLVTIR

Subtype B

**Immunogen** HIV-1 infection, computer prediction

Species (MHC) human (A\*3303)

Assay type Chromium-release assay

Keywords binding affinity, computational epitope predic-

tion

References Hossain et al. 2003

- HLA-A\*3303 is a common HLA allele in east and southeast Asia. Pol, Gag and Nef SF2 proteins were scanned for potential A\*3303 epitopes. 99 potential epitopes were synthesized, and 52/99 bound to A\*3303. Six of these served as peptide-targets for lysis by PBMC from infected individual, and clones derived from 4 of these 6 could lyse HIV-vaccinia infected target cells, indicating proper processing.
- This epitope is one of the 4 that are properly processed.

HXB2 Location Protease (7–15)

**Author Location** Protease

Epitope QRPLVTIKI

Epitope name QI9

Immunogen HIV-1 infection

Species (MHC) human (A\*0101)

**Donor MHC** A\*0101, A\*0205, B\*0702, B\*0801,

Cw\*0701, Cw\*0702

Country Australia.

Assay type Intracellular cytokine staining

**Keywords** HAART, ART, immunotherapy, variant crossrecognition or cross-neutralization, optimal

epitope

References Stratov et al. 2005

 CD8 T-cells targeting epitopes spanning drug resistance induced mutations were detected in 3/25 individuals harboring multidrug-resistant HIV-1. Novel CD8 T-cell responses were detected against epitopes with common protease inhibitor fitness mutations. T-cell immunity to drug-resistant variants was confirmed in SHIV-infected macaques, where CD8 and CD4 immune responses to RT and protease resistance mutations were elicited using peptide-based immunotherapy.

- QRPLVTIKI carries the the L10I protease inhibition mutation and was recognized in a multidrug resistant individual. Response against wild-type epitope qrpIvtiki was detected.
- The 3 people that responded to the drug resistant forms of the virus were among those that had the highest levels of CD4 and CD8 T-cell responses, indicating that they were among the most immunocompetent.

**HXB2 Location** Protease (11–20)

**Author Location** Pol

Epitope VTIKIGGQLK

Epitope name Pol 98

Subtype M

Immunogen vaccine, in vitro stimulation or selection, com-

puter prediction

Vector/Type: DNA, peptide Adjuvant: In-

complete Freund's Adjuvant (IFA)

**Species (MHC)** human, mouse, humanized mouse (A\*1101)

Assay type Cytokine production, T-cell Elispot

**Keywords** subtype comparisons, computational epitope prediction, escape, TCR usage, variant cross-

recognition or cross-neutralization

References McKinney et al. 2004

- This study examined variant recognition of epitopes presented by A\*0201 and A\*1101. Numerous amino acid substitutions can be introduced into epitopes without disrupting their recognition by CTLs, although epitopes with multiple substitutions were less recognized. An algorithm was constructed for prediction of epitopes capable of inducing responses to a great number of variant epitopes.
- A total of 30 variant forms of Pol 98 were identified. 50% of the variant epitopes were recognized by CTLs from mice immunized with the parental form.
- Pol 98 epitope was present in 71% of HIV sequences of many M group subtypes.

HXB2 Location Protease (11–20)

Author Location Pol (91–100)

Epitope VTILIGGQLK

Immunogen HIV-1 infection

**Species (MHC)** human (A3 supertype)

**Keywords** supertype, rate of progression

**References** Propato *et al.* 2001

- B\*0801, Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer ant cross-
  - Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
  - A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
  - This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location Protease (12–20)

Author Location Pol (92–100)
Epitope TIKIGGQLK
Immunogen HIV-1 infection
Species (MHC) human (A3 supertype)
Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location Protease (23–32)

**Author Location** Pol

Epitope LLDTGADDTV

Epitope name L10V

Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV compo-

*nent:* gp140, gp140ΔV3 **Species (MHC)** transgenic mouse (A\*0201)

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted
CTL responses against multiple conserved HIV-1 epitopes in
HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw
virus expressing gp140 with deleted V1, V2, and V3 loops
successfully induces neutralizing antibodies against HIV-1. A
live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination
against HIV and measles.

HXB2 Location Protease (30–38)
Author Location Pol (subtype B)
Epitope DTVLEEMNL

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A\*6802)

Keywords subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi—these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.

- This epitope is conserved among B and D clade viruses.
- The Clade A version of the epitope: DTVLEDINL.
- This epitope was recognized by two different exposed and uninfected prostitutes.
- This epitope was identified by screening 49 HIV-1 peptides with the predicted A\*6802 anchor residue motif x(VT)xxxxxx(VL)

HXB2 Location Protease (30–38)

**Author Location** Pol (subtype A)

**Epitope** DTVLEDINL

Subtype A

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A\*6802)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 IFNγ responses in the cervix—systemic CD8+ T cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

**HXB2 Location** Protease (30–38)

Author Location RT (85–93 subtype D)

**Epitope** DTVLEEWNL

Subtype D

Immunogen

Species (MHC) human (A\*6802)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*6802 epitope.

HXB2 Location Protease (30–38)

Author Location Pol (subtype A)

**Epitope** DTVLEDINL

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (A\*6802)

Keywords HIV exposed persistently seronegative

(HEPS), escape

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- DTVLEDINL was recognized in 3 of the 6 women (ML857, ML1203, and ML1707), and the response was present in the last available sample prior to seroconversion, 3-7 months.
- In each of the three women, 20/20 sequences of the infecting strain had no substitutions in this epitope, all were DTVLEDINL, so there was no evidence for escape.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.

 This epitope was recognized in 3/22 HEPS sex worker controls, ML851, ML1432, and ML1601.

**HXB2 Location** Protease (30–38) **Author Location** Pol (85–93)

Epitope DTVLEDINL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A\*6802)

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-A\*6802 women, 11/12 HEPS and 6/11 HIV-1 infected women recognized this epitope likelihood ratio 4.4, p value 0.08, and HEPS women tended to respond to DTVLEDINL, infected women tended to ETAYFILKL.
- The dominant response to this HLA allele was to this epitope in 10 of the 11/12 HEPS cases, but in only 4 of the 6/11 HIV-1 infected women.
- Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.
- Four epitopes were considered to be "resistant epitopes", as they were preferentially reactive in HEPS women and so may confer resistance, and these were found in three different proteins: A2 ILK(D/E)PVHGV in RT, A\*6802 DTVLEDINL in Protease, B14 DLN(M/T)LN(I/V)V in p24 and B18 FRDYV-DRF(Y/F)K also in p24.
- Subject ML 857 shifted from a A\*6802 DTVLEDINL and B35 (H/N)PDIVIYQY response prior to seroconversion to a B35 PPIPVGDIY and B35 VPLRPMTY response postseroconversion, and the loss of the pre-seroconversion response was not due to sequence variation within these epitopes.
- Subject ML 1203 started with CTL responses to A\*6802 DTVLEDINL and B7 FPVTPQVPLR prior to seroconversion, and upon seroconversion acquired additional responses to A\*6802 ETAYFILKL which became dominant, B7 TPGPGV/IRYPL, B7 IPRRIRQGL, and B7 SPRTLNAWV.
- Subject ML 1707 started with a CTL response to A\*6802 DTVLEDINL prior to seroconversion, and switched to A\*6802 ETAYFILKL and A24 RDYVDRFFKTL post-seroconversion, and the loss of the pre-seroconversion response was not due to sequence variation within the epitope.
- Subject ML 1830 made no detectable response prior to seroconversion, but responded to A\*6802 DTVLEDINL and A\*6802 ETAYFILKL post-seroconversion.

HXB2 Location Protease (30–38)

Author Location Pol

**Epitope** DTVLEDINL

Immunogen HIV-1 infection

Species (MHC) human (A\*6802)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

**HXB2 Location** Protease (30–38)

**Author Location** Pol (87–95)

**Epitope** DTVLEEMNL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A28)

Donor MHC A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** Protease (30–38)

**Author Location** (B consensus)

Epitope DTVLEEMNL

**Epitope name** DL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A68)

**Donor MHC** A31, A68, B07, B70, Cw7, Cw1

Country United States.

Assay type Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

**References** Lichterfeld *et al.* 2004c

 Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.

• 1/9 individuals recognized this epitope.

**HXB2 Location** Protease (34–42) **Author Location** Protease (34–42) Epitope EEINLPGKW Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*44)

Assay type Other

**Keywords** HLA associated polymorphism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- EEINLPGKW was a previously defined B\*44 presented epitope that encompassed an associated polymorphism, EeINLPGKW,in the second position.

**HXB2 Location** Protease (34–42) **Author Location** Protease (34–42)

Epitope EEMNLPGRW

Immunogen

Species (MHC) human (B44)

Keywords optimal epitope References Frahm et al. 2007

**HXB2 Location** Protease (34–42)

**Author Location** Protease

Epitope EEMNLPGRW

**Epitope name** EW9

Immunogen HIV-1 infection Species (MHC) human (B44)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Chromium-release assay, Flow

cytometric T-cell cytokine assay

Keywords epitope processing, supervised treatment interruptions (STI), immunodominance

References Rodriguez et al. 2004

• Protease and integrase are shown to be frequently targeted by CD8 T-cell responses (23% and 68% of 56 HIV+ patients, respectively). Responses tend to cluster in conserved regions of Int, although 1 high conserved region had no responses. CTL frequencies per unit protein length for Pro and Int were similar to other HIV non-structural proteins. Three novel HLA class I-restricted optimal epitopes were found and characterized with fine mapping.

The epitope includes residue M36, which is a known accessory mutation site in individuals treated with PIs.

**HXB2 Location** Protease (45–54) **Author Location** Pol (45–54 IIIB) Epitope KMIGGIGGFI

Epitope name pol45-54

Subtype B Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB

HIV component: Gag-Pol

**Species (MHC)** humanized mouse (A\*0201) Assay type Intracellular cytokine staining

Keywords epitope processing, vaccine-specific epitope characteristics, immunodominance, variant cross-recognition or cross-neutralization, vaccine antigen design

References Singh & Barry 2004

- When A\*0201-C3H/J transgenic mice were immunized with plasmids encoding wild-type gag-pol, codon-optimized (CO) gag-pol, and an expression library vaccine expressing 16 fragments of gag-pol fused with ubiquitin for proteosome targeting (ELI), the ELI vaccine produced up to 10-fold higher CD8 T-cell responses than the other two vaccines. In contrast to the wt and CO vaccines, which tended to augment only immunodominant responses, boosting with the ELI vaccine resulted in many CD8 responses against variant epitopes from different HIV-1 clades, and against drug-resistant variants.
- The drug resistant variant of this epitope, kViVgiggfi, was tested. WT and CO vaccines produced low level CD8+ Tcell responses against the B clade form as well as against drug resistant variant, but the ELI vaccine produced much more intense responses against both the WT and the variant, including after boosting.

**HXB2 Location** Protease (45–54)

Author Location Pol (125-134)

Epitope KMIGGIGGFI Immunogen HIV-1 infection

Species (MHC) human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind three of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

HXB2 Location Protease (57–66) **Author Location** 

Epitope RQYDQILIEI

Epitope name RI10

**Immunogen** 

Species (MHC) human (B13)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B13 epitope.

HXB2 Location Protease (58-66)

**Author Location** Protease

Epitope QYDQIPIEI

Epitope name QI9

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0401)

**Donor MHC** A\*0201, A\*1101, B\*1501, B\*3501

Cw\*0401, Cw\*0701

Country Australia.

Assay type Intracellular cytokine staining

**Keywords** HAART, ART, immunotherapy, variant crossrecognition or cross-neutralization, optimal

epitope

References Stratov et al. 2005

- CD8 T-cells targeting epitopes spanning drug resistance induced mutations were detected in 3/25 individuals harboring multidrug-resistant HIV-1. Novel CD8 T-cell responses were detected against epitopes with common protease inhibitor fitness mutations. T-cell immunity to drug-resistant variants was confirmed in SHIV-infected macaques, where CD8 and CD4 immune responses to RT and protease resistance mutations were elicited using peptide-based immunotherapy.
- QYDQIPIEIW harbors the L63P protease inhibitor mutation, and this created an epitope. The wild-type epitope qydqiLiei was not recognized.
- The 3 people that responded to the drug resistant forms of the virus were among those that had the highest levels of CD4 and CD8 T-cell responses, indicating that they were among the most immunocompetent.

HXB2 Location Protease (68–76)

**Author Location** (C consensus)

Epitope GKKAIGTVL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- GKKAIGTVL is an optimal epitope.

HXB2 Location Protease (68–76)

**Author Location** 

Epitope GKKAIGTVL

Epitope name GL9

Immunogen

Species (MHC) human (B\*1503)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1503 epitope.

HXB2 Location Protease (69–83)

**Author Location** Protease (69–83)

Epitope HKAIGTVLVGPTPVN

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Other

Keywords TCR usage, characterizing CD8+ T cells

References Yang et al. 2005a

- B\*1501, B\*3501, CTL responses were evaluated in identical twins infected with HIV-1 from the same blood source. Targeting of the CTL was similar in the 2 patients, while their TCR profiles were highly dissimilar. It is suggested that CTL targeting is predominately genetically determined, while T-cell generation is a stochastic process; the responding CTLs differ at the TCR molecular level, leaving the viral escape and CTL efficacy unpredictable.
  - HKAIGTVLVGPTPVN was the immunodominant epitope in each twin, but was recognized by T cells with distinctly different TCRs. The epitope was not defined within the epitope. The twins had very different patterns of HIV evolution in this region, with one carrying HKveGsVLiGPTPVN and the other HKAIGaVLiGPTPVN.

**HXB2 Location** Protease (70–77)

**Author Location** 

Epitope KAIGTVLV

Immunogen

Species (MHC) human (B57)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B57 epitope.

**HXB2 Location** Protease (70–77)

**Author Location** Protease

Epitope KAIGTVLV

Epitope name KV8

Subtype B, C

Immunogen HIV-1 infection

**Species (MHC)** human (B63, B57, B58)

**Donor MHC** A23, A30, B42, B57, Cw17

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

 HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.  HLA-B63/57/58 epitope containing the B58 supertype binding motif. Significantly more often recognized by B63-positive subjects than by negative subjects. Optimal epitope was defined in a person who was B57+.

HXB2 Location Protease (75–84)
Author Location Protease (75–84 MN)
Epitope VLVGPTPVNI

Immunogen in vitro stimulation or selection

**Species (MHC)** human (A\*0201) **Keywords** binding affinity **References** Konya *et al.* 1997

- Peptide predicted to be reactive based on HLA-A\*0201 binding motif.
- Peptide could stimulate CTL in PBMC from 5/6 seronegative donors.
- Peptide located in a highly conserved region of protease.
- Both 9-mer and 10-mer could stimulate CTL: VLVGPTPVNI and LVGPTPVNI.
- Binding affinity to A\*0201 was measured,  $C_{1/2 \text{ max}} \mu M = 6$  for 10-mer, 3 for 9-mer.
- MAL variant of Pr(75-84 MN), with substitutions V77, G78, and P79, gave reduced binding and CTL recognition.

HXB2 Location Protease (75–84)

Author Location Protease (175–184 MN)

Epitope VLVGPTPVNI

**Subtype** B **Immunogen** vaccine

Vector/Type: DNA, polyepitope Strain: B clade MN HIV component: gp120, Protease, RT Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) humanized mouse (A\*0201) Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance, immunotherapy

References Isaguliants et al. 2004

- Immunization of HLA-A\*0201-transgenic mice with synthetic genes encoding clusters of human A\*0201 CTL epitopes located at the sites of drug resistance mutations, induced RTspecific cellular responses indicating the immunogenicity of these constructs. This vaccine strategy may be a first step towards a therapeutic vaccine against drug-resistant strains.
- This was one of five HLA-A\*0201 epitopes from the RT or protease proteins that was included in the polyepitope vaccine. When the transgenic HLA A\*0202 mice were vaccinated with the polyepitope construct or with a mixture of RT peptides, a sustained low level CD8+ T-cell gamma IFN response was observed, in contrast to when an intact RT gene was used for vaccination.

HXB2 Location Protease (75–84)
Author Location Pol (75–84 IIIB)
Epitope VLVGPTPVNI
Epitope name pol75-84
Subtype B
Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB HIV component: Gag-Pol

Species (MHC) humanized mouse (A\*0201)

Assay type Intracellular cytokine staining

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance, variant cross-recognition or cross-neutralization, vaccine antigen design

References Singh & Barry 2004

- When A\*0201-C3H/J transgenic mice were immunized with plasmids encoding wild-type gag-pol, codon-optimized (CO) gag-pol, and an expression library vaccine expressing 16 fragments of gag-pol fused with ubiquitin for proteosome targeting (ELI), the ELI vaccine produced up to 10-fold higher CD8 T-cell responses than the other two vaccines. In contrast to the wt and CO vaccines, which tended to augment only immunodominant responses, boosting with the ELI vaccine resulted in many CD8 responses against variant epitopes from different HIV-1 clades, and against drug-resistant variants.
- The drug resistant variant of this epitope,vlvgptpTnV, was tested. WT and CO vaccines produced low level CD8+ T-cell responses against the B clade form as well as against drug resistant variant, but the ELI vaccine produced much more intense responses against both the WT and the variant, including after boosting.

**HXB2 Location** Protease (76–84)

**Author Location** Pol (163–)

Epitope LVGPTPVNI

Epitope name Pol-163

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** binding affinity, subtype comparisons, supertype, computational epitope prediction

References Altfeld et al. 2001c

- HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.
- Three additional previously described HLA-A2 epitopes were added to the set of 20, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- LVGPTPVNI binds to 4/5 HLA-A2 supertype alleles: A\*0201, A\*0202, A\*0206 (highest affinity) and A\*6802, but not A\*0203.
- 1/22 individuals with chronic HIV-1 infection recognized this epitope by ELISPOT.
- 0/12 acutely infected individuals recognized this epitope.

HXB2 Location Protease (76–84) Author Location Protease (76–84) Epitope LVGPTPVNI

**Epitope name** LI9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a A\*0201 epitope.

**HXB2 Location** Protease (76–84)

**Author Location** Protease

Epitope LVGPTPANI

Immunogen HIV-1 infection

Species (MHC) human, macaque (A\*0205)

**Donor MHC** A\*0101, A\*0205, B\*0702, B\*0801,

Cw\*0701, Cw\*0702

Country Australia.

Assay type Intracellular cytokine staining

Keywords HAART, ART, immunotherapy, variant crossrecognition or cross-neutralization

References Stratov et al. 2005

- CD8 T-cells targeting epitopes spanning drug resistance induced mutations were detected in 3/25 individuals harboring multidrug-resistant HIV-1. Novel CD8 T-cell responses were detected against epitopes with common protease inhibitor fitness mutations. T-cell immunity to drug-resistant variants was confirmed in SHIV-infected macaques, where CD8 and CD4 immune responses to RT and protease resistance mutations were elicited using peptide-based immunotherapy.
- A response against the peptide harboring the protease drug resistance mutation V82A, LVGPTPANI, was detected in one individual, but the wildtype epitope was not recognized, lygptpVni. This epitope response was not fine-mapped, and is based on analogy to a previously described A2 epitope.
- Other drug resistant variants were not recognized: V82T and
- The 3 people that responded to the drug resistant forms of the virus were among those that had the highest levels of CD4 and CD8 T-cell responses, indicating that they were among the most immunocompetent.

**HXB2 Location** Protease (76–84)

**Author Location** Protease (76–84 HXB2)

Epitope LVGPTPVNI Epitope name PR82V

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type Intracellular cytokine staining, Chromiumrelease assay

Keywords HAART, ART, escape References Karlsson et al. 2003

- This epitope contains two positions that are commonly associated with protease inhibitor escape, lvgptpAni (V82A) and lvgptpvnV (I84V). 29 HIV-1 infected patients (15 were HLA-A2+) with a history of protease inhibitor failure were screened for mutations within the protease gene and CD8+ T cells recognition of the wt and V82A variant peptides. CTL pressure alone, despite high functional avidity, did not drive the V82A substitution. Suprisingly V82A was found more frequently among HLA-A2- individuals (10/14) than HLA-A2+ (7/15), despite the mutation conferring not only drug resistance but CTL escape.
- 8/15 HLA-A2+ patients carried had a Val at position 82; 7/8 of these recognized the WT peptide, but only 3/8 could also recognize V82A.

• 7/15 had the V82A substitution; 2/7 recognized the wt and the V82A mutation, 1/7 recognized only the peptide with the V82A substitution.

**HXB2 Location** Protease (76–84)

**Author Location** Protease (76–84)

Epitope LVGPTPVNI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Canada.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

assay

Keywords HAART, ART, escape, immunotherapy, variant cross-recognition or cross-neutralization

References Mason et al. 2004

- · Accumulation of specific antiretroviral drug-resistance mutations in Pol gene was shown to sustain and even enhance the antigenicity and immunogenicity of HIV-1 CTL epitopes in this region. Several different patterns of cross-reactivity and selective recognition of wild-type and variant epitopes were found.
- LVGPTPVNv variant is detected due to appearence of I84V resistance mutation. Three patients receiving PIs had viral sequences obtained, and two had the I84V mutation. EliSpot reactivity to this epitope in either form was evident in these patients, showing drug resistnace can persist coincident with an active CTL response.

HXB2 Location Protease (76–84)

**Author Location** Protease (76–84)

Epitope LVGPTPVNI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

• The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized during chronic infection.

HXB2 Location Protease (76–84)

**Author Location** Protease (76–84)

Epitope LVGPTPVNI

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Canada.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

assay, Flow cytometric T-cell cytokine assay

Keywords variant cross-recognition or crossneutralization, characterizing CD8+ T

cells, mimics

References Mason et al. 2005

- CTL responses against the human IP-30 signal peptide sequence LLDVPTAAV were shown to be elicited by stimulation of PBMCs from HIV-1 infected individuals with HIV protease peptide 76-84, LVGPTPVNI. In vitro stimulation with HIV PR 76-84 or the IP-30 signal peptide was shown to activate a comparable population of cross-reactive effector cells. None of the peptides activated CTL in non-HIV-infected individuals. IP-30 signal peptide was shown to have lower avidity T-cell interactions than the HIV peptide.
- As a control, responses to A2-restricted HIV epitopes ALVE-ICTEM, EELRQHLLRW, and LSPRTLNAW were shown not to give IP-30 responses.

HXB2 Location Protease (76–84)

Author Location Pol (156–164)

Epitope LVGPTPVNI

Immunogen HIV-1 infection

Species (MHC) human (A2 supertype)

Keywords supertype, rate of progression
References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind three of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

HXB2 Location Protease (76–85)
Author Location (C consensus)
Epitope LVGPTPVNII
Subtype C
Immunogen HIV-1 infection
Species (MHC) human (A\*0205)

**Country** South Africa. **Assay type** CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- LVGPTPVNII is an optimal epitope.

HXB2 Location Protease (79–89)

**Author Location** Protease

Epitope PTPVNIIGRNL

Subtype B, C

Immunogen HIV-1 infection Species (MHC) human (B63)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- Putative HLA-B63/57/58 epitope containing the B58 supertype binding motif. Significantly more often recognized by B63positive subjects than by negative subjects, trend towards being more often recognized in those with B57/B58.

HXB2 Location Protease (80–90)
Author Location (C consensus)
Epitope TPVNIIGRNML
Subtype C
Immunogen HIV-1 infection

Species (MHC) human (B\*8101)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- TPVNIIGRNML is an optimal epitope.

**HXB2 Location** Protease (80–90)

**Author Location** 

Epitope TPVNIIGRNML

Epitope name TL11

Immunogen

Species (MHC) human (B81)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B81 epitope.

## II-B-11 Protease-RT CTL/CD8+ epitopes

**HXB2 Location** Protease-RT (95–5) **Author Location** Gag (175–184)

Epitope CTLNFPISPI

Immunogen HIV-1 infection

**Species (MHC)** human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- The epitope starts in Protease and ends in RT.
- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.

- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind all five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802)

HXB2 Location Protease-RT (96–5)
Author Location Pol (176–184)
Epitope TLNFPISPI
Immunogen HIV-1 infection
Species (MHC) human (A2 supertype)
Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind three of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

HXB2 Location Protease-RT Author Location Protease-RT

**Epitope** 

Immunogen SHIV infection, vaccine

*Vector/Type:* peptide *HIV component:* Protease, RT

Species (MHC) macaque

Assay type Intracellular cytokine staining

**Keywords** HAART, ART, vaccine-specific epitope char-

acteristics, vaccine-induced epitopes, escape, immunotherapy

References Stratov et al. 2005

- CD8 T-cells targeting epitopes spanning drug resistance induced mutations were detected in 3/25 individuals harboring multidrug-resistant HIV-1. Novel CD8 T-cell responses were detected against epitopes with common protease inhibitor fitness mutations. T-cell immunity to drug-resistant variants was confirmed in SHIV-infected macaques, where CD8 and CD4 immune responses to RT and protease resistance mutations were elicited using peptide-based immunotherapy.
- The SHIV infected macaques that responded best to the peptide vaccine were those that did not yet have progressive disease. Thus peptide immunotherapy for multidrug resistance has the best hope of success if given to those who are not yet fully immunocompromised.

## II-B-12 RT CTL/CD8 + epitopes

HXB2 Location RT (1-16)

**Author Location** (C consensus)

Epitope PISPIETVPVKLKPGM

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*3910)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location RT (3-12)

**Author Location RT (LAI)** 

Epitope SPIETVPVKL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2, B61)

References van der Burg et al. 1997

- Recognized by CTL from a long-term survivor, EILKEPVGHGV was also recognized.
- Highly conserved across clades.

HXB2 Location RT (3-12)

**Author Location** (C consensus)

Epitope SPIETVPVKL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*8101)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the P2 residue of SPIETVPVKL are associated with the presence of the HLA presenting molecule in the host.

HXB2 Location RT (3-12)

**Author Location** Pol

Epitope SPIETVPVKL

Immunogen

Species (MHC) human (B7)

References De Groot et al. 2001

The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.

- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes could stimulate IFNy production in an ELISPOT assay.
- SPIETVPVKL was newly identified as HLA-B7 epitope in this study, it had been previously shown to be presented by HLA-A2 and B61.

HXB2 Location RT (3-12)

**Author Location** Pol

Epitope SPIETVPVKL

Epitope name 1307

Subtype multiple

Immunogen HIV-1 infection

**Species (MHC)** human (B7, A2, B8, B61)

Donor MHC A03, A24, B07, B38, Cw07, Cw12/13; A29, Author Location (C consensus)

A30, B08, B44, Cw07, Cw16

Country United States. Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion, cross-presentation by different HLA

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for SPIETVPVKL: 12%. Promiscuous epitope binding to A02, B07, B08 and B61.

HXB2 Location RT (5-12)

**Author Location** RT (5–12)

Epitope IETVPVKL

**Immunogen** HIV-1 infection

Species (MHC) human (B\*4001)

**Keywords** optimal epitope

References Frahm et al. 2007

HXB2 Location RT (5-29)

**Author Location** RT (160–184 HXB2)

Epitope IETVPVKLKPGMDGPKVKQWPLTEE

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Walker et al. 1989

• One of five epitopes defined for RT-specific CTL clones in this

HXB2 Location RT (14-23)

**Author Location** Pol

Epitope PGMDGPKVKQ

Epitope name 1276

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A11)

Donor MHC A11, A68, B42, B45, Cw16, Cw17

Country United States.

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope prediction, cross-presentation by different HLA

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for PGMDGPKVKQ:52% Promiscuous epitope binding to A11 or A68, previously published

HXB2 Location RT (15-32)

Epitope GMDGPKVKQWPLTEEKIK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4202)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location RT (18-26)

Author Location RT (185–193 LAI)

Epitope GPKVKQWPL

Subtype B

**Immunogen** 

Species (MHC) human (B\*0801)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*0801 epitope.

HXB2 Location RT (18-26)

**Author Location** RT (18-26)

Epitope GPKVKQWPL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Meier et al. 1995; Menendez-Arias et al. 1998

- HIV proteins with mutations in this epitope allowed transactive inhibition of specific CTL-mediated lysis.
- Article reviewed in Menendez-Arias et al. [1998], with a discussion of antagonism.

HXB2 Location RT (18-26)

**Author Location** RT (173–181)

**Epitope** GPKVKQWPL

Immunogen

Species (MHC) human (B8)

References Goulder et al. 1997g; Menendez-Arias et al. 1998

• Included in a study of the B8 binding motif.

cussion of antagonism.

HXB2 Location RT (18-26)

**Author Location** RT (185–193 LAI)

Epitope GPKVKQWPL

Subtype B

Immunogen

Species (MHC) human (B8)

References Sutton et al. 1993

· Predicted epitope based on B8-binding motifs, from larger peptide IETVPVKLKPGMDGPKVKQWPLTEE.

HXB2 Location RT (18–26)

**Author Location** RT (185–193 LAI)

**Epitope** GPKVKQWPL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Klenerman et al. 1995; Menendez-Arias et al.

- Naturally occurring antagonist GPRVKQWPL found in viral PBMC DNA and RNA.
- Article reviewed in Menendez-Arias et al. [1998] with a discussion of antagonism.

HXB2 Location RT (18-26)

Author Location RT (18–26)

Epitope GPKVKQWPL

Immunogen in vitro stimulation or selection

Species (MHC) human (B8)

Keywords dendritic cells

References Zarling et al. 1999

- This study compares the ability of macrophages and dendritic cells to stimulate primary responses in CD8+ lymphocytes isolated from HLA-appropriate HIV-uninfected donors using peptide-pulsed APC – the dendritic cells performed better as APC for the stimulation of primary responses.
- · Strong CTL responses were elicited by the epitopes DRFYK-TLRA and GEIYKRWII when presented by either immature or mature dendritic cells - macrophages were not able to prime a CTL response against DRFYKTLRA.
- A weak response to KLTPLCVSL was stimulated using macrophages as the APC.
- · No detectable response was observed for the following previously-defined HIV epitopes: KIRLRPGGK, ILKEPVHGV, IRLRPGGK, GPKVKQWPL.

HXB2 Location RT (18-26)

Author Location RT (185-193)

Epitope GPKVKQWPL

**Epitope name** GPK

Immunogen HIV-1 infection Species (MHC) human (B8)

Keywords HAART, ART, supervised treatment inter-

ruptions (STI), immunodominance, escape,

acute/early infection

References Oxenius et al. 2000

- Article reviewed in Menendez-Arias et al. [1998], with a diswith sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load - three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
  - Two of the 7/8 study subjects that were HLA B8+ recognized this epitope.
  - Patient SC2 (HLA A1, B7/8, Cw0701/0702, DR4/53, DQ7) had CTL responsiveness against epitopes FLKEKGGL, GP-KVKQWPL, and GEIYKRWII peptides - FLKEKGGL tetramer staining steadily declined and at day 1340 the FLKEKGGL stained cells were no longer detected and the escape mutant FLKENGGI was found in 8/10 clones.
  - Patient SC11(HLA A1, B8, Cw0201, DR3/11, DR52, DQ2/7) started therapy early, remained on therapy for 40 days, then reinitiated HAART at day 640 had a CTL response to FLKEKGGL, GPKVKQWPL, and GEIYKRWII throughout and received a benefit from the early limited course therapy.

HXB2 Location RT (18-26)

**Author Location** Pol

Epitope GPKVKQWPL

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords HAART, ART

References Seth et al. 2001

• CTL responses were studied by tetramer staining in 41 patients with combination therapy - activated CD8+ T-cells decline as the viral load drops in response to therapy, but the overall level of antigen-specific cells capable of differentiating into effectors stays constant and new epitopes may be recognized.

HXB2 Location RT (18-26)

**Author Location** RT (185–193 SF2)

Epitope GPKVKQWPL

**Immunogen** HIV-1 infection

Species (MHC) human (B8)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B8+ individuals that had a CTL response to this epitope broken down by group: 0/3 group 1, 0/3 group 2, and 2/2 group 3.

HXB2 Location RT (18-26)

**Author Location** Pol (171–180)

 ${\bf Epitope}\ {\sf GPKVKQWPL}$ 

Subtype A, B, C, D

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B8)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

• GPKVKQWPL is cross-reactive for clades A, B, C, and D.

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

HXB2 Location RT (18-26)

**Author Location** RT (18–26)

Epitope GPKVKQWPL

Immunogen HIV-1 infection

Species (MHC) human (B8)

**References** Day *et al.* 2001

• B8-restricted CTL accounted for about 1/3 of the total CTL response in one individual.

HXB2 Location RT (18-26)

Author Location RT

Epitope GPKVKQWPL

Epitope name GPK

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords HAART, ART, supervised treatment interrup-

tions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius et al. [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there was no correlation between CTL responses with viral rebound rates, plateau viral loads, or clearance rates.

HXB2 Location RT (18-26)

Author Location Pol (171–180)

Epitope GPKVKQWPL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ, CD8 T-cell Elispot

granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- One of the patients responded to this peptide with GzB producing cells, while two different patients responded with IFN-gamma producing cells.

HXB2 Location RT (18-26)

Author Location (B consensus)

Epitope GPKVKQWPL

Epitope name GL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

Donor MHC A25, A32, B08, B14, Cw7, Cw8

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
   activation in dying target cells, it was shown that the subset of
   HIV-1-specific CD8+ T cells secreting both IFN-gamma and
   TNF-alpha exhibit stronger cytotoxic activity than those secret ing only IFN-gamma. These cells also exhibited stronger in tracellular perforin expression. No association between HIV-1 specific CD8+ T-cell maturation phenotypes and intracellular
   perforin expression was found.
- 1/9 individuals recognized this epitope.

HXB2 Location RT (18-26)

**Author Location** Pol (173–181)

Epitope GPKVKQWPL

Epitope name GL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A\*01, A\*11, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay

Keywords escape, variant cross-recognition or cross-

neutralization, optimal epitope

References Koibuchi et al. 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- The GL9 variant GPrVKQWPL was essentially the only form of the epitope detected over a 5-year period in this person. Elispot reactions were roughly equivalent between the autologous form and the B clade consensus form, GPKVKQWPL. A single variant was observed in 1/8 clones at the 5-year time point, GPrVKOgPL.

HXB2 Location RT (18–27)

**Author Location** Pol

Epitope GPKVKQWPLT

Immunogen

Species (MHC) human (B7, B8)

References De Groot et al. 2001

- gram Conservatrix to identify conservered regions of HIV that might serve as epitopes.
- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes were identified that could stimulate IFN production in an ELISPOT assay.
- GPKVKOWPLT was confirmed as a previously identified HLA-B8 epitope, and newly identified as an HLA-B7 epitope in this study.

HXB2 Location RT (18-27)

**Author Location** Pol

Epitope GPKVKQWPLT

Epitope name 1293

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (B7, B8)

**Donor MHC** A03, A24, B07, B38, Cw07, Cw12/13; A29,

A30, B08, B44, Cw07, Cw16

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope prediction, cross-presentation by different HLA

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC
- Estimated binding probability for GPKVKQWPLT: 27% Promiscuous epitope binding to B07 and B08.

HXB2 Location RT (33-41)

**Author Location** RT (33–41 LAI)

**Epitope** ALVEICTEM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0201 epitope.

HXB2 Location RT (33-41)

Author Location RT (33-41 LAI)

**Epitope** ALVEICTEL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords binding affinity, computational epitope predic-

tion

References Samri et al. 2000

• This epitope contains the mutation M41L, a mutation induced by nucleosidee reverse transcriptase inhibitors.

- The program Epimatrix was used in conjunction with the prothe mutated peptide after zidovudine treatment, but not the wild-type peptide - the mutation M41L gave an increased A2 binding score (http://bimas.dcrt.nih.gov/molbio/hla\_bind) compared to the wildtype RT sequence.
  - Three additional A\*0201 individuals and one B27 individual did not respond to this epitope before or after treatment.
  - M41L occurred at anchor positions p2 and p9 in several computer predicted RT epitopes (33-41, 32-41, and 40-49) (http//bimas.dcrt.nih.gov/molbio/hla\_bind), and increased the predicted binding affinity for 6 HLA molecules (B2705, B5102, C3, A0201, B2705 and B3901)

HXB2 Location RT (33-41)

Author Location RT (33-41 MN)

Epitope ALVEICTEM

Subtype B

Immunogen vaccine

Vector/Type: DNA, polyepitope Strain: B clade MN HIV component: gp120, Protease, RT Adjuvant: Incomplete Freund's Adju-

vant (IFA)

**Species (MHC)** humanized mouse (A\*0201)

Assay type CD8 T-cell Elispot - IFNγ

Keywords epitope processing, vaccine-specific epitope characteristics, immunodominance, im-

munotherapy

References Isaguliants et al. 2004

- Immunization of HLA-A\*0201-transgenic mice with synthetic genes encoding clusters of human A\*0201 CTL epitopes located at the sites of drug resistance mutations, induced RTspecific cellular responses indicating the immunogenicity of these constructs. This vaccine strategy may be a first step towards a therapeutic vaccine against drug-resistant strains.
- This was one of five HLA-A\*0201 epitopes from the RT or protease proteins that was included in the polyepitope vaccine. When the transgenic HLA A\*0202 mice were vaccinated with the polyepitope construct or with a mixture of RT peptides, a sustained low level CD8+ T-cell gamma IFN response was observed, in contrast to when an intact RT gene was used for vaccination.

HXB2 Location RT (33-41)

**Author Location** Pol (132–140 IIIB)

**Epitope** ALVEICTEM

Epitope name pol132-140

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB

HIV component: Gag-Pol

Species (MHC) humanized mouse (A\*0201)

Assay type Intracellular cytokine staining

Keywords subtype comparisons, vaccine-specific epitope characteristics, immunodominance, vari-

ant cross-recognition or cross-neutralization,

vaccine antigen design

References Singh & Barry 2004

- When A\*0201-C3H/J transgenic mice were immunized with plasmids encoding wild-type gag-pol, codon-optimized (CO) gag-pol, and an expression library vaccine expressing 16 fragments of gag-pol fused with ubiquitin for proteosome targeting (ELI), the ELI vaccine produced up to 10-fold higher CD8 T-cell responses than the other two vaccines. In contrast to the wt and CO vaccines, which tended to augment only immunodominant responses, boosting with the ELI vaccine resulted in many CD8 responses against variant epitopes from different HIV-1 clades, and against drug-resistant variants.
- Different variants of this epitope from different clades were tested. WT and CO vaccines produced low level CD8+ T-cell responses against the B clade form as well as against variants from other clades, but the ELI vaccine produced much more intense responses against the B clade and all variants tested, including after boosting. The variants were: clade A, alTDictem; clade C, atTAicEem; and clade D, alIeicSem.

HXB2 Location RT (33-41)

**Author Location** Pol

Epitope ALVEICTEM

Epitope name A9M

Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV compo-

nent: gp140, gp140 $\Delta$ V3

Species (MHC) transgenic mouse (A\*0201)

Assay type Chromium-release assay Fl

**Assay type** Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted
CTL responses against multiple conserved HIV-1 epitopes in
HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw
virus expressing gp140 with deleted V1, V2, and V3 loops
successfully induces neutralizing antibodies against HIV-1. A
live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination
against HIV and measles.

HXB2 Location RT (33–41)

**Author Location** RT (33–41)

Epitope ALVEICTEM

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Haas et al. 1998

- Of 98 patients in cross-sectional analysis, 78% had CTL against pol – RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)
- New clusters of epitopes were defined utilizing different HLA molecules.

HXB2 Location RT (33-41)

Author Location RT (33-41)

**Epitope** ALVEICTEM

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- SLYNTVATL was the dominant A2 epitope recognized in patients with chronic infection, except for Subject 11841 who recognized 5/8 epitopes and who had a dominant A-2 response to ALVEICTEM.

HXB2 Location RT (33-41)

Author Location RT (33-41)

**Epitope** ALVEICTEM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

 The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized in chronic infection, and even then was recognized infrequently.

HXB2 Location RT (33-41)

**Author Location** RT (33–41)

Epitope ALVEICTEM

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Canada.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords mimics

References Mason et al. 2005

- CTL responses against the human IP-30 signal peptide sequence LLDVPTAAV were shown to be elicited by stimulation of PBMCs from HIV-1 infected individuals with HIV protease peptide 76-84, LVGPTPVNI. In vitro stimulation with HIV PR 76-84 or the IP-30 signal peptide was shown to activate a comparable population of cross-reactive effector cells. None of the peptides activated CTL in non-HIV-infected individuals. IP-30 signal peptide was shown to have lower avidity T-cell interactions than the HIV peptide.
- As a control, responses to A2-restricted HIV epitopes ALVE-ICTEM, EELRQHLLRW, and LSPRTLNAW were shown not to give IP-30 responses.

HXB2 Location RT (33-41)

**Author Location** RT (33–41)

**Epitope** ALVEICTEM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2, A3)

Country Canada.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords HAART, ART, immunotherapy, variant cross-

recognition or cross-neutralization

References Mason et al. 2004

- Accumulation of specific antiretroviral drug-resistance mutations in Pol gene was shown to sustain and even enhance the antigenicity and immunogenicity of HIV-1 CTL epitopes in this region. Several different patterns of cross-reactivity and selective recognition of wild-type and variant epitopes were found.
- ALVEICTEI variant is detected due to appearence of M41L resistance mutation. The M41L variant peptide was almost always preferentially recognized by CTLs from patients undergoing antiretroviral therapy.

HXB2 Location RT (33-43)

Author Location RT (33-43)

**Epitope** ALVEICTEMEK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

References Haas et al. 1998

- Of 98 patients in cross-sectional analysis, 78% had CTL against pol – RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)
- New clusters of epitopes were defined utilizing different HLA molecules.
- C. Brander notes that this is an A\*0301 epitope in the 1999 database, G. Haas, pers. comm.

HXB2 Location RT (33-43)

Author Location RT (33-43)

**Epitope** ALVEICTEMEK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0301 epitope.

HXB2 Location RT (33-43)

Author Location RT (33-43)

**Epitope** ALVEICTEMEK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- All patients recognized at least 1 A3 epitope, up to 8 A3 epi- Epitope noted in a review by Menendez-Arias et al. [1998] to topes, but none was clearly dominant.

HXB2 Location RT (33-43)

Author Location RT Pol (188-198)

**Epitope** ALVICTEMEK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong T-helper cell responses. Only patients starting with moderately high viral load (VL) were able to reduce the VL set point. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow
- Less than 2 of 14 patients recognized this epitope.

HXB2 Location RT (38-52)

Author Location RT (203-209)

**Epitope** CTEMEKEGKISKIGP

Immunogen vaccine

Vector/Type: Salmonella HIV component:

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Burnett et al. 2000

· A live attenuated bacterial vaccine, Salmonella SL3261pHART, with an inserted HIV epitope in the Lpp-OmpA-HIV fusion protein, induced a specific CTL response in BALB/c mice (<15% lysis assayed by Cr-release of target cells)

HXB2 Location RT (38-52)

**Author Location** RT (205–219 BRU)

Epitope CTEMEKEGKISKIGP

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: RT

**Species (MHC)** mouse (H-2<sup>k</sup>)

Keywords review

References De Groot et al. 1991; Menendez-Arias et al.

1998

- Murine and human helper and CTL epitope.
- Epitope noted in a review by Menendez-Arias et al. [1998] to be located in the "fingers" domain of RT and is a helper and CTL epitope.

HXB2 Location RT (38-52)

**Author Location** RT (205–219)

**Epitope** CTEMEKEGKISKIGP

Immunogen HIV-1 infection

Species (MHC) human (broad)

Keywords review

References Hosmalin et al. 1990; Menendez-Arias et al. 1998

- Murine and human helper and CTL epitope.
- be located in the "fingers" domain of RT and is a helper and CTL epitope.

HXB2 Location RT (39-47)

**Author Location RT** 

**Epitope** TEMEKEGKI

Immunogen

**Species (MHC)** mouse (H-2K<sup>k</sup>)

References Leggatt et al. 1998

- Epitope variants were examined for CTL response in concert with H-2K<sup>k</sup> MHC class I binding all of the following combinations were observed: (i) two single mutations which did not alone abrogated CTL activity did abrogate activity when combined, (ii) loss of recognition of a single substitution could be restored by an additional substitution, and (iii) sometimes there was recognition of two single substitutions as well as the combination of those substitutions.
- 2E and 9I are anchor residues for H-2K<sup>k</sup> if you have M in the third position, it enhances H-2K<sup>k</sup> binding 10-fold, but polymorphism at this site is important for the overall conformation of the peptide and can influence T cell recognition.

HXB2 Location RT (39-47)

Author Location RT (206-214)

**Epitope TEMEAEGKI** 

Immunogen in vitro stimulation or selection

Species (MHC) mouse

Keywords TCR usage

References Leggatt et al. 1997

- Ala-substituted nonamer-peptide used to test a non-radioactive assay for murine CTL recognition of peptide-MHC class I complexes.
- The new assay is CTL adherence assay (CAA), and is based on the discovery that CTL develop adhesive properties upon TCR triggering.
- Substitutions in TEMEAEGKI that reduce cytolytic activity were correctly detected by CAA.

HXB2 Location RT (42-50)

**Author Location** RT (42–50 LAI)

**Epitope** EKEGKISKI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*5101 epitope.

HXB2 Location RT (42-50)

**Author Location** RT (42–50 HXB2)

Epitope EKEGKISKI

Epitope name EI9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

• T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.

HXB2 Location RT (42-50)

Author Location RT (42-50 LAI)

Epitope EKEGKISKI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B51)

References Haas et al. 1998

- Of 98 patients in cross-sectional analysis, 78% had CTL against pol – RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)
- New clusters of epitopes were defined utilizing different HLA molecules.

HXB2 Location RT (42–50)

**Author Location** RT (42–50)

**Epitope** EKEGKISKI

Immunogen HIV-1 infection

Species (MHC) human (B51)

**Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

HXB2 Location RT (55-72)

**Author Location** (C consensus)

Epitope PYNTPVFAIKKKDSTKWR

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location RT (57-65)

Author Location RT

Epitope NTPVFAIKK

Immunogen HIV-1 infection, vaccine

Species (MHC) human (A\*6801, A3 superfamily)

Country Australia.

Assay type Intracellular cytokine staining

**Keywords** HAART, ART, immunotherapy, variant crossrecognition or cross-neutralization, optimal

epitope

References Stratov et al. 2005

- CD8 T-cells targeting epitopes spanning drug resistance induced mutations were detected in 3/25 individuals harboring multidrug-resistant HIV-1. Novel CD8 T-cell responses were detected against epitopes with common protease inhibitor fitness mutations. T-cell immunity to drug-resistant variants was confirmed in SHIV-infected macaques, where CD8 and CD4 immune responses to RT and protease resistance mutations were elicited using peptide-based immunotherapy.
- The immune response to this peptide was cross-reactive for both the wild type and RT drug resistance mutation K65R, and NTPVFAIKK and ntpvfaikR stimulated CD8 T-cell responses with equal efficiency. The C-terminal R or K is required for a full response; NTPVFAIK stimulated a much weaker response.
- The 3 people that responded to the drug resistant forms of the virus were among those that had the highest levels of CD4 and CD8 T-cell responses, indicating that they were among the most immunocompetent.

HXB2 Location RT (57–65)

**Author Location** Pol (236–244)

**Epitope** NTPVFAIKK **Immunogen** HIV-1 infection

**Species (MHC)** human (A3 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location RT (57-66)

Author Location Pol

Epitope NTPVFAIKKK

Epitope name 1274

Subtype multiple

Immunogen HIV-1 infection

**Species (MHC)** chimpanzee, goat, baboon (A11, A68, B8)

**Donor MHC** A01, A68, B15, B40, Cw03; A25, A68, B18,

B27

**Country** United States. **Assay type** T-cell Elispot

Keywords binding affinity, supertype, computational epi-

tope prediction, immunodominance, cross-

presentation by different HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for NTPVFAIKKK:53%. Epitope binds to A11 and A68 supertype, and is immunodominant.

HXB2 Location RT (73-82)

Author Location RT (73-82)

Epitope KLVDFRELNK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

HXB2 Location RT (73-82)

**Author Location** RT (73–82 LAI)

**Epitope** KLVDFRELNK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**References** Samri *et al.* 2000

- This epitope contains the mutation L74V, a frequent mutation induced by nucleoside reverse transcriptase inhibitors.
- The wild-type, but not the mutated peptide, was recognized before and after zidovudine treatment in A3-restricted patients 252#0 and 252#4.
- Mutation L74V affects the p2 anchor position in RT epitopes and was predicted to reduce binding to A3 (http://bimas.dcrt.nih.gov/molbio/hla bind)

HXB2 Location RT (73-82)

Author Location RT (228-237)

Epitope KLVDFRELNK

Epitope name A3-KK10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

 CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.

- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 3/7 individuals began to have detectable responses to this epitope after STI.

HXB2 Location RT (73–82) Author Location RT (73–82) Epitope KLVDFRELNK Epitope name A3-KK10 Pol Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3) Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords supervised treatment interruptions (STI), es-

cape, early treatment, superinfection

References Altfeld et al. 2002a

 An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response, but in this case the epitope did not vary.

HXB2 Location RT (73-82)

**Author Location** Pol

**Epitope** KLVDFRELNK

Subtype multiple

Immunogen HIV-1 infection Species (MHC) human (A3)

**Donor MHC** A03, A23, B49, B57; A02, A03, B08, B51, Cw01, Cw07; A03, A11, B14, B05, Cw08

**Country** United States. **Assay type** T-cell Elispot

**Keywords** binding affinity, computational epitope prediction

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for KLVDFRELNK: 36%.

HXB2 Location RT (73–82)
Author Location (B consensus)
Epitope KLVDFRELNK
Epitope name KK10
Subtype B
Immunogen HIV-1 infection

Species (MHC) human (A3)
Donor MHC A03, B07, Cw7
Country United States.

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

HXB2 Location RT (87–104)

**Author Location** (C consensus)

Epitope FWEVQLGIPHPAGLKKKK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location RT (93-101)

**Author Location** (LAI)

**Epitope** GIPHPAGLK

Subtype B

Immunogen

Species (MHC) human (A\*0301)

Keywords optimal epitope

References Altfeld 2000; Frahm et al. 2007

**HXB2 Location** RT (93–101)

Author Location RT (248-257)

Epitope GIPHPAGLK

Epitope name A3-GK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

 CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.

- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 1/7 individuals began to have detectable responses to this epitope after STI.

**HXB2 Location** RT (93–101) **Author Location** RT (93–101)

**Epitope** GIPHPAGLK **Epitope name** A3-GK9 Pol

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3) Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

 An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response, but in this case the epitope did not vary.

HXB2 Location RT (93-101)

**Author Location** Pol

**Epitope** GIPHPAGLK

Epitope name 1337

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A03, A23, B49, B57

**Country** United States. **Assay type** T-cell Elispot

**Keywords** binding affinity, computational epitope predic-

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for GIPHPAGLK: 20%.

HXB2 Location RT (93–101)

**Author Location** (B consensus)

Epitope GIPHPAGLK

Epitope name GK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A03, B07, Cw7

Country United States.

Assay type Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

HXB2 Location RT (93–102)

**Author Location** Pol (240–249 93TH253 subtype CRF01)

Epitope GIPHPAGLKK Epitope name P248-257 Subtype CRF01\_AE

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was weakly reactive in the HEPS study subject 265 who was HLA A2/A11 and after a second stimulation in vitro gave a strong response in HEPS study subject 128 who was HLA A11/A33.

HXB2 Location RT (93-102)

**Author Location** Pol (240–249 93TH253 subtype CRF01)

Epitope GIPHPAGLKK
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A11)

**Keywords** subtype comparisons

References Bond et al. 2001

- HLA-A11 CRF01 (called subtype E in Bond *et al.*) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This is one of the new A11 epitopes identified through the streamlined EpiMatrix method, and 2/8 tested FSWs recognized it.

• This epitope was highly conserved in other subtypes, and exact **Author Location** RT (262–270 IIIB) matches were common.

HXB2 Location RT (98-113) **Author Location** RT (252–266) Epitope AGLKKKKSVTVLDVGD Immunogen HIV-1 infection Species (MHC) human (Cw4) References Bernard et al. 1998

- This study focuses on six rare long-term survivor HIV-infected people who were infected for many years without exhibiting immune dysregulation - such immunologically normal HIVinfected (INHI) cases occur at a frequency between 0.1 and 1% in the infected population.
- No direct CTL were found in any of the six INHIs, but above background CTLp activity was founded in 3/6 INHIs.

HXB2 Location RT (98-113)

Author Location Pol (254–264 BH10, LAI)

Epitope AGLKKKKSVTVLDVGD

Immunogen HIV-1 infection

Species (MHC) human

- References Maksiutov et al. 2002
- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is GLKKKKSVTVL) has similarity with the CD166 antigen (activated leukocyte-cell adhesion molecule), fragment GLKKRESLTLI.

**HXB2 Location** RT (102–118) **Author Location** (C consensus)

Epitope KKKSVTVLDVGDAYFSV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0401)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (103–117) Author Location RT (257–251) Epitope KKSVTVLDVGDAYFS Immunogen HIV-1 infection Species (MHC) human (Cw4) References Bernard et al. 1998

- This study focuses on six rare long-term survivor HIV-infected people who were infected for many years without exhibiting immune.
- No direct CTL were found in any of the six INHIs, but above background CTLp activity was founded in 3/6 INHIs.

**HXB2 Location** RT (107–115)

Epitope TVLDVGDAY

**Immunogen** 

**Species (MHC)** (B\*3501)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*3501 epitope.

**HXB2 Location** RT (107–115)

Author Location Pol (262-270)

Epitope TVLDVGDAY

Epitope name TY9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*3501)

Donor MHC A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Chromium-release assay

Keywords escape, acute/early infection

References Bansal et al. 2005

- · Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- The response to this epitope was not evident until month 20, and increased over time.

**HXB2 Location** RT (107–115)

**Author Location** RT (262–270 IIIB)

Epitope TVLDVGDAY Immunogen HIV-1 infection Species (MHC) human (B35)

Keywords review, responses in children, mother-to-

infant transmission

References Menendez-Arias et al. 1998; Wilson et al.

1996

- Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.
- TVLDMGDAC is a naturally occurring variant that is less reactive.
- Menendez-Arias et al. [1998], in a review, notes that this epitope includes a catalytic residue (Asp-110) in the active site of RT.

**HXB2 Location** RT (107–115)

**Author Location** Pol (262–270 IIIB)

**Epitope** TVLDVGDAY Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords responses in children, mother-to-infant transmission, escape

References Wilson et al. 1999a

• This study describes maternal CTL responses in the context of mother-to-infant transmission.

- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- An additional variant that gave a positive CTL response: TVLDMGDAC.

**HXB2 Location** RT (107–115) Author Location Pol (262–270) **Epitope** TVLDVGDAY Immunogen HIV-1 infection Species (MHC) human (B35) References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** RT (107–115) **Author Location** RT (262–270 SF2) **Epitope** TVLDVGDAY Immunogen HIV-1 infection Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- · The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B35+ individuals that had a CTL response to this epitope broken down by group: 1/2 group 1, 0/2 group 2, and 0/1 group 3.

**HXB2 Location** RT (107–115) **Author Location** 

Epitope TVLDVGDAY Epitope name Pol-TY9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35) References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA B35, 8/21 (38%) recognized this epitope.

**HXB2 Location** RT (107–115)

Author Location Pol

**Epitope** TVLDVGDAY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35) **Donor MHC** A3, A11, B35, B51

Keywords mother-to-infant transmission

References Sabbaj et al. 2002

- IFNgamma T-cell responses in breast milk of 5 HIV-1 infected women from the US and 6 from Zambia were tested with using Elispot. 11/11 women responded to Gag, 8/11 responded to Pol, 7/11 women to Nef, and 2/5 women to Env peptide pools. These responses were shown to be primarily due to CD8+ T-cells in one woman, and another woman had cytolytic responses measured by Cr-release.
- T-cells in breast milk from a volunteer who was HLA A3, A11, B35, B51 induced IFNgamma after stimulation with either of two overlapping peptides that carry known B35 epitope TVLDVGDAY.
- The frequencies of responses in the two compartments differed, and 2/4 women that responded to epitopes in Nef 101-205 and Pol 601-710 showed responses in breast milk but no detectable responses in peripheral blood cells.

**HXB2 Location** RT (107-115)

Author Location RT (107-115)

**Epitope** TVLDVGDAY

Subtype AG

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country Canada.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords HAART, ART, immunotherapy, variant crossrecognition or cross-neutralization

References Mason et al. 2004

- · Accumulation of specific antiretroviral drug-resistance mutations in Pol gene was shown to sustain and even enhance the antigenicity and immunogenicity of HIV-1 CTL epitopes in this region. Several different patterns of cross-reactivity and selective recognition of wild-type and variant epitopes were
- TiLDVGDAY, TVLDVGDAf and TiLDVGDAf variants are detected due to appearence of V108I and Y115F resistance mutations. Complete cross-reactivity of wild-type and variant peptides was observed.

HXB2 Location RT (107-115)

**Author Location** RT Pol (262–270)

**Epitope** TVLDVGDIY

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFNγ, Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/9 patients recognized this epitope.

**HXB2 Location** RT (108–118) Author Location RT (267-277) Epitope VLDVGDAYFSV

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

References van der Burg et al. 1996

- High dissociation rate, but immunogenic in primary CTL induction after repeated stimulations with peptide.
- CTL generated by in vitro stimulation of PBMC derived from uninfected individual.

HXB2 Location RT (108-118)

**Author Location** Pol

**Epitope** VLDVGDAYFSV

Epitope name V11V Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV component: gp140, gp140ΔV3

**Species (MHC)** transgenic mouse (A\*0201)

Assay type Chromium-release assay, Flow cytometric Tcell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted
CTL responses against multiple conserved HIV-1 epitopes in
HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw
virus expressing gp140 with deleted V1, V2, and V3 loops
successfully induces neutralizing antibodies against HIV-1. A
live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination
against HIV and measles.

HXB2 Location RT (108–118)

Author Location RT (267–277)

Epitope VLDVGDAYFSV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords dendritic cells

References Kundu et al. 1998b

- Allogeneic dendritic cells (DCs) were obtained from HLAidentical siblings, pulsed with rgp160 MN or A2-restricted HIV-1 epitope peptides, and infused monthly into six HIVinfected patients.
- 1/6 showed increased env-specific CTL and increased lymphoproliferative responses, 2/6 showed increase only in proliferative responses, and 3/6 showed no change pulsed DCs were well tolerated.
- VLDVGDAYFSV is a conserved HLA-A2 epitope included in this study – 4/6 patients had this sequence as their HIV direct sequence, but only one of these had a detectable CTL response – the other two had the sequences EEDVGDAYFSV and ELDVGDAYFSV and no detectable CTL response.

HXB2 Location RT (108-118)

Author Location RT (267–277)

Epitope VLDVGDAYFSV

Immunogen in vitro stimulation or selection

Species (MHC) human (A2)

References van der Burg et al. 1995

- Binds HLA-A\*0201 CTL generated by in vitro stimulation of PBMC from an HIV negative donor.
- VLDVGDAYFSV is in a functional domain.

**HXB2 Location** RT (108–118)

**Author Location** RT Pol (263–273)

**Epitope** VLDVGDAYFSV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 19 patients recognized this epitope.

HXB2 Location RT (108-118)

**Author Location** Pol (263–273)

Epitope VLDVGDAYFSV

Immunogen HIV-1 infection

Species (MHC) human (A2, A\*0201)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** RT (108–122)

**Author Location** RT (257–251)

Epitope VLDVGDAYFSVPLDE

Immunogen HIV-1 infection

Species (MHC) human (Cw4)

**References** Bernard *et al.* 1998

- This study focuses on six rare long-term survivor HIV-infected people who were infected for many years without exhibiting immune dysregulation – such immunologically normal HIVinfected (INHI) cases occur at a frequency between 0.1 and 1% in the infected population.
- No direct CTL were found in any of the six INHIs, but above background CTLp activity was founded in 3/6 INHIs.

**HXB2 Location** RT (113–120)

**Author Location** Pol (268–275 SF2)

Epitope DAYFSVPL

Immunogen HIV-1 infection

Species (MHC) human (B\*5101, A24)

Keywords subtype comparisons, rate of progression

References Tomiyama et al. 1999

• HLA-B27, -B51, and -B57 are associated with slow progression to AIDS, while HLA -B35, -B8, -B24 are associated with a rapid progression to AIDS (Nat. Med. 2:405, 1996;Lancet 22:1187, 1986;Hum Immunol 22:73, 1988;Hum Immunol 44:156, 1995)

- 15% of Japanese populations carry HLA-B51 while HLA-B27 and -B57 are detected in less than 0.3%
- Of the 172 HIV-1 peptides with HLA-B\*5101 anchor residues, 33 bound to HLA-B\*5101, seven of these peptides were reactive with CTL from 3 B\*5101 positive individuals, and six were properly processed.
- Four of the six epitopes were highly conserved among B subtype sequences, DAYFSVPL is conserved.

**HXB2 Location** RT (113–120)

Author Location RT (113–120)

Epitope DAYFSVPL

Immunogen HIV-1 infection

Species (MHC) human (B51)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

**HXB2 Location** RT (116–124)

Author Location (C consensus)

Epitope FSVPLDEDF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*35)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the E7 residue of FSVPLDEDF are associated with the presence of the HLA presenting molecule in the host.
- FSVPLDEDF not optimized.

**HXB2 Location** RT (116-124)

Author Location (C consensus)

Epitope FSVPLDEDF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

**References** Kiepiela *et al.* 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- FSVPLDEDF is an optimal epitope.

**HXB2 Location** RT (116–124)

Author Location (C consensus)

**Epitope** FSVPLDEDF

Subtype C

**Immunogen** HIV-1 infection

Species (MHC) human (B\*5703)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- FSVPLDEDF is an optimal epitope.

**HXB2 Location** RT (116–135)

Author Location Pol (271–290)

Epitope FSVPLDEDFRKYTAFTIPSI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons

References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** RT (117–126)

Author Location Pol (264–273 93TH253 subtype CRF01)

**Epitope** SVPLDESFRK **Epitope name** P272-281

Subtype CRF01\_AE

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.

• This epitope after a second stimulation in vitro gave a strong • A K to E substitution at position 5 abrogates specific lysis, and response in HEPS study subject 128 who was HLA A11/A33.

**HXB2 Location** RT (117-126)

Author Location Pol (264–273 93TH253 subtype CRF01)

**Epitope** SVPLDESFRK Subtype CRF01\_AE Immunogen HIV-1 infection Species (MHC) human (A11)

Keywords subtype comparisons References Bond et al. 2001

- HLA-A11 CRF01 (called subtype E in Bond et al.) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This is one of the new A11 epitopes identified through the streamlined EpiMatrix method, and 3/8 tested FSWs recognized it.
- This epitope was only conserved in CRF01, and subtype A and B, and exact matches were uncommon.

**HXB2 Location** RT (118–127)

Author Location (C consensus)

**Epitope** VPLDEDFRKY

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*35)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** RT (118–127)

**Author Location** RT (273–282 SF2)

Epitope VPLDKDFRKY

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

Keywords review

References Menendez-Arias et al. 1998; Tomiyama et al.

- A CTL clone responsive to this epitope was obtained.
- 4/7 B35-positive individuals had a CTL response to this epitope.

- reduces binding to B\*3501.
- Menendez-Arias et al. [1998], in a review, notes that a Glu to Lys (E to K) change abrogates CTL activity, but that both VPLDEDFRKY and VPLDKDFRKY can serve as HLA-B35 epitopes, so the change must alter T cell receptor binding residues in this epitope may be important for polymerase activ-

**HXB2 Location** RT (118–127)

**Author Location** RT (273–282 IIIB)

Epitope VPLDEDFRKY

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*3501 epitope.

**HXB2 Location** RT (118–127)

Author Location Pol (273–282)

Epitope VPLDKDFRKY

**Immunogen** HIV-1 infection

Species (MHC) human (B\*3501)

References Tomiyama et al. 2000a

- CD8+ T-cells that bound one of six HIV-specific B\*3501epitope tetramers did not express CD28 or CD45A.
- A significant increase in CD28-CD45RA- cells and a decrease of CD28+CD45RA+ cells was observed in chronically HIV-1infected individuals relative to healthy individuals.
- CD28-CD45RA- cells are likely to be effector cells and have high levels of perforin in their cytoplasm.
- The mean percentage of total CD28- CD8+ cells in chronically infected HIV-1-infected patients was 76.6% in comparison to HIV-1-uninfected individuals (40.6%)

HXB2 Location RT (118-127)

**Author Location (SF2)** 

Epitope VPLDEDFRKY

Epitope name HIV-B3501-SF2-4

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

References Tomiyama et al. 2000b

• B\*3501 VPLDEDFRKY tetramer binding did not inhibit CTL activity of a clone that react with both HLA-B\*3501 than HLA-B\*5101 presentation of the epitope IPLTEEAEL.

**HXB2 Location** RT (118–127)

Author Location RT (118-127)

**Epitope** VPLDEDFRKY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

**Donor MHC** A\*2301, B\*3501, B\*1503, Cw2, Cw7

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

**HXB2 Location** RT (118–127) Author Location Pol (273–282) **Epitope** VPLDEDFRKY

Epitope name VY10 Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

**Donor MHC** A\*0201, A\*0301, B\*3501, B\*51, Cw\*04, Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Chromium-release assay

Keywords escape, acute/early infection

References Bansal et al. 2005

- · Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- The response to this epitope was not detected until month 25, and increased over time.

**HXB2 Location** RT (118–127)

**Author Location** RT (273–282 IIIB)

**Epitope** VPLDEDFRKY

Immunogen HIV-1 infection

**Species (MHC)** human (B\*3501, B35)

References Shiga et al. 1996

• Binds HLA-B\*3501.

HXB2 Location RT (118-127)

**Author Location (SF2)** 

Epitope VPLDKDFRKY Immunogen HIV-1 infection Species (MHC) human (B35)

**Keywords** binding affinity, rate of progression, escape References Kawana et al. 1999

- HLA B35 is associated with rapid disease progression.
- The sequences of 9 previously described HIV-1 B35 CTL epitopes were obtained in 10 HLA B35+ and 19 HLA B35- individuals.
- 3/9 CTL epitopes had substitutions that were more common in B35+ individuals than in B35- individuals – only one of these reduced the binding of the peptide to B35 and was shown to be an escape mutation.
- -E--- was found in 8/10 of the B35+ individuals, and three of the B35- individuals – the D –> E substituted peptide had similar binding affinity to B35 and was equally susceptible to a CTL clone.

HXB2 Location RT (118-127)

**Author Location** RT (273–282 IIIB)

**Epitope** VPLDEDFRKY

Immunogen HIV-1 infection

Species (MHC) human (B35)

**Keywords** subtype comparisons

References Sipsas et al. 1997

- HIV IIIB proteins were used to define the range of CTL epitopes recognized by three lab workers accidentally infected with HIV-1 IIIB.
- · VPLDKDFRKY, a variant found in HIV MN, was not recog-
- · VPHDEDFRKY, a variant found in HIV YU2, was not recog-
- · This epitope was type-specific and conserved in only one other B subtype sequence.

HXB2 Location RT (118-127)

**Author Location** RT (273–282 SF2)

Epitope VPLDEDFRKY

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- · Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B35+ individuals that had a CTL response to this epitope broken down by group: 1/2 group 1, 0/2 group 2, and 1/1 group 3.

**HXB2 Location** RT (118-127)

**Author Location** 

Epitope VPLDEDFRKY

Epitope name Pol-VY10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35) References Sabbaj *et al.* 2003

 Among HIV+ individuals who carried HLA B35, 5/21 (24%) recognized this epitope.

**HXB2 Location** RT (118–127)

**Author Location** RT Pol (273–282)

Epitope VPLDEDFRKY Immunogen HIV-1 infection Species (MHC) human (B35)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/9 patients recognized this epitope.

HXB2 Location RT (126–135)
Author Location RT (293–302 HXB)
Epitope KYTAFTIPSI
Immunogen HIV-1 infection
Species (MHC) human (A2)
Keywords HAART, ART

References Shankar *et al.* 1998
• A novel CTL clone was defined with a panel of recombinant

by a patient who was HIV-seropositive for 6 years and had not received any antiretroviral therapy.
There is evidence that some CTL epitopes are poorly presented on the surface of infected cells, but this RT epitope was recognized.

vaccinia-RT-infected B-LCL target cells using PBMCs donated

 There is evidence that some CTL epitopes are poorly presented on the surface of infected cells, but this RT epitope was recognized as effectively on HIV-infected cells as on peptide-pulsed targets.

HXB2 Location RT (127–135)

**Author Location** (C consensus)

**Epitope** YTAFTIPSI

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*0205)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- YTAFTIPSI is an optimal epitope.

**HXB2 Location** RT (127–135)

Author Location Pol (316-)

Epitope YTAFTIPSI

**Epitope name** Pol-316

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** binding affinity, subtype comparisons, supertype, computational epitope prediction

References Altfeld et al. 2001c

- HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.
- Three additional previously described HLA-A2 epitopes were added to the set of 20, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- 2/22 individuals with chronic HIV-1 infection recognized this epitope in ELISPOT.
- 0/12 acutely infected individuals recognized this epitope.
- YTAFTIPSI binds to five HLA-A2 supertype alleles: A\*0201, A\*0202,A\*0203, A\*0206 and A\*6802 (highest affinity)

**HXB2 Location** RT (127–135)

Author Location RT (127-135)

**Epitope YTAFTIPSV** 

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords optimal epitope

**References** Frahm *et al.* 2007

**HXB2 Location** RT (127–135)

Author Location RT (127-135)

**Epitope** YTAFTIPSI

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2)

Country United States.

Country Officer States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

 The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized during chronic infection.

**HXB2 Location** RT (127-135)

**Author Location** Pol (306–314)

Epitope YTAFTIPSI

Immunogen HIV-1 infection

Species (MHC) human (A2 supertype)

**Keywords** supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind all five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802)

 $\textbf{HXB2 Location} \ \ RT \ (128\text{--}135)$ 

Author Location Pol (283–290)

**Epitope** TAFTIPSI

Epitope name TI8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

 $\textbf{Donor MHC} \ A*0201, \ A*0301, \ B*3501, \ B*51, \ Cw*04,$ 

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Chromium-release assay

**Keywords** escape, acute/early infection, characterizing

CD8+ T cells

References Bansal et al. 2005

- Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- The response to this epitope was weak and sporadic.

**HXB2 Location** RT (128-135)

**Author Location** 

Epitope TAFTIPSI

**Epitope name** Pol-TI8

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (A\*0217, B\*5101)

**Donor MHC** 01RCH46: A\*0201, A\*0217, B\*0801,

B\*4002, Cw\*0303, Cw\*0701

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.

- Patient 01RCH46 was Hispanic, on HAART, and had a viral load of 21000 and CD4 count of 623 she also recognized GELDRWEKI, p17(11-19), HLA B\*4002, and KETINEEAA p24(70-78), HLA B\*4002.
- Among HIV+ individuals who carried HLA A\*02, 7/36 (19%) recognized this epitope, two of which also carried B\*5101 which can also restrict this epitope.

**HXB2 Location** RT (128–135)

Author Location RT (295–302 IIIB)

**Epitope TAFTIPSI** 

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

 $\textbf{Keywords} \ \ \text{optimal epitope}$ 

References Frahm et al. 2007

• C. Brander notes this is a B\*5101 epitope.

**HXB2 Location** RT (128-135)

**Author Location** Pol (283–290 SF2)

Epitope TAFTIPSI

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

**Keywords** subtype comparisons, rate of progression

References Tomiyama et al. 1999

- HLA-B27, -B51, and -B57 are associated with slow progression to AIDS, while HLA-B35, -B8, -B24 are associated with a rapid progression to AIDS (Nat. Med. 2:405, 1996;Lancet 22:1187, 1986;Hum Immunol 22:73, 1988;Hum Immunol 44:156, 1995)
- 15% of Japanese populations carry HLA-B51 while HLA-B27 and -B57 are detected in less than 0.3%
- Of the 172 HIV-1 peptides with HLA-B\*5101 anchor residues, 33 bound to HLA-B\*5101, seven of these peptides were reactive with CTL from 3 B\*5101 positive individuals, and six were properly processed.
- Four of the six epitopes were highly conserved among B subtype sequences, but TAFTIPSI is somewhat variable.

**HXB2 Location** RT (128–135)

Author Location RT (295–302)

Epitope TAFTIPSI

Epitope name P5

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

Keywords HAART, ART, escape

References Samri et al. 2000

• The epitope TAFTIPSI was recognized by patient 246#1 in a study of the effects of therapy escape mutations on CTL recognition.

**HXB2 Location** RT (128–135)

**Author Location** RT (128–135 IIIB)

**Epitope TAFTIPSI** 

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B\*5101)

Keywords epitope processing, escape

References Moore et al. 2002b

- HIV polymorphisms in the RT protein were examined relation to HLA alleles found in a population of 473 participants in the Western Australian HIV Cohort Study. 64 significant associations between polymorphisms at particular positions and HLA alleles were detected, for HLA-B7, -B12, -B35 and -B15. Fifteen of these were in positions with known epitopes, 4 in anchor residues, 11 in other positions. Six additional polymorphic sites associated with particular HLA molecules flanked known epitopes and may relate to processing.
- TAFTIPSI was one of two epitopes characterized in detail. C-terminal I135x substitutions were associated with people who carried HLA-B5 39/40 (98%) of HLA-B\*5101 individuals had substitutions in this position, while only 127/431 (29%) who did not have HLA-B\*5101 did. The predominant substitution was kytaftipsT, and this mutation is predicted to abrogate binding to HLA-B\*5101.

HXB2 Location RT (128–135) Author Location RT (128–135 HXB2)

**Epitope TAFTIPSI** 

Epitope name TI8 Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101, Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** escape, immune evasion, optimal epitope, HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- Last position (8) of the epitope had potentially experienced positive selection. TAFTIPSt, TAFTIPSr and TAFTIPSv escape variants were found.

**HXB2 Location** RT (128–135)

**Author Location** RT (295–302 IIIB)

**Epitope** TAFTIPSI

Immunogen HIV-1 infection

Species (MHC) human (B51)

Keywords review

**References** Menendez-Arias *et al.* 1998; Sipsas *et al.* 1997

- HIV IIIB proteins were used to define the range of CTL epitopes recognized by 3 lab workers accidentally infected with HIV-1 IIIB.
- TAFTIPST, a variant found in HIV-1 CAM1, was also recognized but 100-fold more peptide was needed.
- TAFTIPSV, a variant found in HIV-1 VE1RT, was also recognized, but 10-fold more peptide was needed.
- TVFTIPSI, a variant found in HIV-1 MANC, was also recognized

Menendez-Arias et al. [1998], in a review, notes that this epitope includes a region near the active site of RT – the substitution of the position two conservative change from A to V decreases CTL recognition.

**HXB2 Location** RT (128–135)

Author Location RT (295-302)

Epitope TAFTIPSI

Immunogen HIV-1 infection

Species (MHC) human (B51)

 $Keywords \ {\it immunodominance}$ 

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- Three of the four individuals that responded to SLYNTVATL recognized additional HIV epitopes, and all three were also HLA B51 and recognized this epitope as well as other epitopes.

**HXB2 Location** RT (128–135)

Author Location RT (295–302)

Epitope TAFTIPSI

**Epitope name** TAF

Immunogen HIV-1 infection

Species (MHC) human (B51)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- None of the 8 study subjects recognized this epitope but none were HLA B51+

**HXB2 Location** RT (128–135)

Author Location RT (295–302 LAI)

Epitope TAFTIPSI

Epitope name P5

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B51)

Keywords HAART, ART

References Mollet et al. 2000

 A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.

In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened – eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+PBL – but with continued viral suppression, HIV-specific responses diminished.

 Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

**HXB2 Location** RT (128-135)

Author Location Pol

**Epitope TAFTIPSI** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B51)

**Donor MHC** A3, A11, B35, B51

Keywords mother-to-infant transmission

References Sabbaj et al. 2002

- IFNgamma T-cell responses in breast milk of 5 HIV-1 infected women from the US and 6 from Zambia were tested with using Elispot. 11/11 women responded to Gag, 8/11 responded to Pol, 7/11 women to Nef, and 2/5 women to Env peptide pools. These responses were shown to be primarily due to CD8+ T-cells in one woman, and another woman had cytolytic responses measured by Cr-release.
- T-cells in breast milk from a volunteer who was HLA A3, A11, B35, B51 induced IFNgamma after stimulation with either of two overlapping peptides that carry known B51 epitope TAFTIPSI.
- The frequencies of responses in the two compartments differed, and 2/4 women that responded to epitopes in Nef 101-205 and Pol 601-710 showed responses in breast milk but no detectable responses in peripheral blood cells.

**HXB2 Location** RT (128–135)

**Author Location** RT (128–135)

**Epitope TAFTIPSI** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B51)

**Donor MHC** A\*0201, A11, B51, B61, Cw2, Cw\*14

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

**HXB2 Location** RT (128-135)

Author Location RT (128-135)

Epitope TAFTIPSI

Immunogen HIV-1 infection

Species (MHC) human (B51)

**Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 5 epitopes from this individual that varied over time where the internal mutations became fixed; a dramatic decrease in CTL activity against the wild type epitope was observed as the mutation arose. The taftipsT variant residue found at 47 months postseroconversion.

**HXB2 Location** RT (128-135)

**Author Location** Pol

Epitope TAFTIPSI

Epitope name TI8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B51)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 8, taftipsT, was found in the most polymorphic residue in the epitope. This was shared between clades B and C.

**HXB2 Location** RT (128–135)

Author Location Pol (295–302)

**Epitope TAFTIPSI** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B51)

**Donor MHC** A2, A31, B51, B58w4

Country United States.

Assay type Intracellular cytokine staining, Flow cytomet-

ric T-cell cytokine assay

Keywords HAART, ART, escape, variant cross-

recognition or cross-neutralization, character-

izing CD8+ T cells

References Casazza et al. 2005

- Evidence of continued viral evolution during HAART therapy despite low viral load was found in 1/5 patients studied. This epitope provided the best evidence for apparent immune escape during HAART.
- Prior to the initiation of therapy, taftipsT variant was found in 24/24 clones. At week 14 of therapy, this variant was completely replaced with taftipsI. By week 19, a complete replacement occurred again, this time to taftipsM. The change at nucleotide level suggests a stepwise progression from ACA to ATA to ATG.
- The taftipsT and taftipsM variants had lower avidity than the taftipsI variant, but this wasn't evident at saturating conditions; only careful titrations revealed the difference. HLA-B51 stabilization studies revealed the increased stabilization with the taftipsI form. Also, CD3 down-regulation was larger in response to taftipsI.
- The viral shift to the taftipsM variant during HAART was predicted to have minimal or undetectable effect on drug sensitivity.

**HXB2 Location** RT (130-144) Author Location RT (130-144)

**Epitope FTIPSINNETPGIRY** 

Immunogen HIV-1 infection

Species (MHC) human (A25)

Assay type Chromium-release assay

Keywords assay standardization/improvement

References Lubong et al. 2004

• Using IL7 or IL15 in culturing of HIV-1-specific CTL clones was inferior to using IL-2 alone; the addition of these cytokines to IL-2 did not show any advantage. Neither proliferation, survival, nor lytic capacity of HIV-1-specific CTLs was significantly enhanced by addition of IL7 or IL15.

**HXB2 Location** RT (136–144)

Author Location RT (136–144 HXB2)

Epitope NNETPGVRY

Epitope name NY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1801)

**Donor MHC** A\*0201, A\*2501,

B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assav type CD8 T-cell Elispot - IFNγ

Kevwords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- NNETPGiRY, NsETPGVRY, NNEiPGiRY, NNEiPGiRY, NNgTPGVRY and NNEvPGiRY escape variants were found.

**HXB2 Location** RT (137-146)

Author Location (C consensus)

Epitope NETPGIRYQY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*18)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the E2 residue of NETPGIRYQY are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** RT (137–146)

**Author Location** 

Epitope NETPGIRYQY

Epitope name NY10

Immunogen

Species (MHC) human (B18)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B18 epitope.

**HXB2 Location** RT (142–149)

**Author Location** (C consensus)

Epitope IRYQYNVL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1401)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- IRYOYNVL is an optimal epitope.

HXB2 Location RT (142-149)

**Author Location** 

Epitope IRYQYNVL

Epitope name IL9

**Immunogen** 

Species (MHC) human (B\*1401)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1401 epitope.

**HXB2 Location** RT (149–159)

**Author Location** (C consensus)

Epitope LPQGWKGSPAI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*3910)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- · LPQGWKGSPAI is an optimal epitope.

**HXB2 Location** RT (151–159)

Author Location Pol (306–314 SF2)

Epitope QGWKGSPAI Immunogen HIV-1 infection Species (MHC) human (B\*5101)

Keywords subtype comparisons, rate of progression

References Tomiyama et al. 1999

- HLA-B27, -B51, and -B57 are associated with slow progression to AIDS.
- 15% of Japanese populations carry HLA-B51 while HLA-B27 and -B57 are detected in less than 0.3%
- Of the 172 HIV-1 peptides with HLA-B\*5101 anchor residues, 33 bound to HLA-B\*5101, seven of these peptides were reactive with CTL from 3 B\*5101 positive individuals, and six were properly processed.
- Four of the six epitopes were highly conserved among B subtype sequences, QGWKGSPAI is conserved.

**HXB2 Location** RT (151–168)

**Author Location** RT (151–168 HXB2)

Epitope QGWKGSPAIFQSSMTKIL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.

 Responses to this peptide were detected in 19% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** RT (153–165)

Author Location RT (308-320)

Epitope WKGSPAIFQSSMT

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Keywords** responses in children, mother-to-infant transmission

References Brander & Walker 1995

• Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.

**HXB2 Location** RT (153–165)

Author Location Pol (308–320)

Epitope WKGSPAIFQSSMT

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** RT (153–167)

Author Location RT (SF2)

Epitope WKGSPAIFQSSMTKI

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2001a

- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- RT peptides SQIYPGIKVRQLCKL and WKG-SPAIFQSSMTKI were recognized.

**HXB2 Location** RT (156–164)

**Author Location** RT (156–164)

**Epitope** SPAIFQSSM

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*07)

Assay type Other

Keywords HLA associated polymorphism

**References** Boutwell & Essex 2007

• All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.

• SPAIFQSSM was a previously defined B\*07 presented epitope that encompassed a B\*0702-associated polymorphism, SPAIFQsSM,in the seventh position.

HXB2 Location RT (156–164) Author Location RT (311–319 SF2) Epitope SPAIFQSSM Immunogen HIV-1 infection Species (MHC) human (B\*3501)

Keywords review

**References** Menendez-Arias *et al.* 1998; Tomiyama *et al.* 1997

- A CTL clone responsive to this epitope was obtained.
- Only 1/7 B35-positive individuals had a CTL response to this epitope.
- Menendez-Arias et al. [1998], in a review, notes that this epitope is near the active site of RT.

HXB2 Location RT (156–164)
Author Location (C consensus)
Epitope SPAIFQSSM
Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*4202) Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- SPAIFQSSM is an optimal epitope.

HXB2 Location RT (156–164) Author Location (C consensus) Epitope SPAIFQSSM

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*8101, B\*0702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** cross-presentation by different HLA, characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** RT (156–164)

**Author Location** RT (311–319 SF2)

Epitope SPAIFQSSM

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords review

References Menendez-Arias et al. 1998; Shiga et al. 1996

- Binds HLA-B\*3501.
- Menendez-Arias et al. [1998], in a review, notes that this epitope includes catalytic residues in the active site of RT.

 $\textbf{HXB2 Location} \ \ RT\ (156\text{--}164)$ 

Author Location Pol (311–319)

Epitope SPAIFQSSM

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location RT (156–164)
Author Location RT Pol (311–319)
Epitope SPAIFQSSM

Immunogen HIV-1 infection Species (MHC) human (B35)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses.
   HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/9 patients recognized this epitope.

**HXB2 Location** RT (156–164)

Author Location Pol (156–164 HXB2)

Epitope SPAIFQSSM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords rate of progression, immunodominance

References Hay et al. 1999

- CTL response to IPRRIRQGL was the immunodominant response in a rapid progressor there was a subdominant response to SPAIFQSSM in Pol, and interestingly, no response to commonly immunodominant HLA A\*0201 epitope SLYNT-VATL, although this individual was HLA A\*0201.
- The individual showed a strong initial CTL response at the time of the initial drop in viremia, but it was quickly lost, although memory cells persisted.
- Despite the initial narrow response to two epitopes, no other CTL responses developed.
- No HIV-specific lymphoproliferative responses were detected in this patient, and neutralizing antibody response was weak.

Variants of this epitopes were observed *in vivo* (spaifqCsm, spSifqssm), but the binding motifs for B7 were preserved (P2, and C-term aromatic or hydrophobic)
 One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two

**HXB2 Location** RT (156-164)

Author Location Pol

Epitope SPAIFQSSM Immunogen HIV-1 infection Species (MHC) human (B7)

 $\boldsymbol{Keywords}$  rate of progression, acute/early infection

References Islam et al. 2001

- Subject 053i was followed longitudinally from acute infection through death, and had rapid progression to AIDS.
- This individual had a dominant response to IPRRIRQGL with strong in vivo activated responses and in vitro stimulated memory responses and a subdominant response to SPAIFQSSM – during the course of disease progression (4 Years), the functional CTL responses were lost and no sequence variation occurred with in both epitopes, but CTL clones specific for IPRIRQGL persisted throughout.

**HXB2 Location** RT (156–164)

Author Location RT (323–331 SF2)

Epitope SPAIFQSSM Immunogen HIV-1 infection Species (MHC) human (B7)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B7+ individuals that had a CTL response to this epitope broken down by group: 0/4 group 1, 0/3 group 2, and 1/1 group 3.

**HXB2 Location** RT (156–164)

Author Location RT (156-164)

Epitope SPAIFQSSM

Epitope name B7-SM9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

 CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.

- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 1/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 3/4 individuals had detectable responses to this epitope after STI.

**HXB2 Location** RT (156–164)

Author Location RT (156-164)

Epitope SPAIFQSSM

**Epitope name** B7-SM9 Pol

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

 An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response; this epitope did not vary.

HXB2 Location RT (156-164)

**Author Location** 

Epitope SPAIFQSSM

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** assay standardization/improvement, epitope

processing

References Draenert et al. 2004a

- 96% of optimally defined epitopes have one of only nine amino acids serving as the C-terminal anchor position. Seven amino acids are never found in this position and four are only present in 4% of cases. CD8 T-cell response to an epitope is shown to be best detected when the epitope is situated at the C-terminal end of a longer peptide, and authors suggest that Elispot reagents would be better designed if peptides ended on known C-terminal anchors.
- SPAIFQSSM is suggested to be the optimal epitope instead of SPAIFQSSMT.

**HXB2 Location** RT (156–164)

Author Location (B consensus)

Epitope SPAIFQSSM

Epitope name SM9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A03, B07, Cw7

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** RT (156–164)

**Author Location** Pol

**Epitope** SPAIFQSSM

Epitope name SM9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 7, spaifqCsm, was found in the most polymorphic residue in the epitope. This was shared between clades B and C.

**HXB2 Location** RT (156–164)

**Author Location** Pol (312–320)

**Epitope** SPAIFQSSM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assav type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** RT (156–164)

**Author Location** 

Epitope SPAIFQSSM

Epitope name SM9

Immunogen

Species (MHC) human (B7)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B07 epitope.

**HXB2 Location** RT (156–165)

**Author Location** RT (311–319 LAI)

Epitope SPAIFQSSMT

Epitope name P4

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords HAART, ART, escape

References Samri et al. 2000

- This epitope contains the mutation P157S which can be induced by nucleosidee reverse transcriptase inhibitors.
- It was recognized by patient 252#0 in a study of the effects of therapy escape mutations on CTL recognition.

**HXB2 Location** RT (156–165)

**Author Location** RT (311–319 SF2)

**Epitope** SPAIFQSSMT

Immunogen

Species (MHC) human (B7)

Keywords review

**References** Brander & Walker 1997; Menendez-Arias *et al.* 1998

- Pers. comm. from C. Hey and D. Ruhl to C. Brander and B. Walker.
- Menendez-Arias et al. [1998], in a review, notes that this epitope includes catalytic residues in the active site of RT.

**HXB2 Location** RT (156–165)

**Author Location** RT (311–319 SF2)

Epitope SPAIFQSSMT

Epitope name P4

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords HAART, ART

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+PBL but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

**HXB2 Location** RT (156–165)

Author Location Pol

Epitope SPAIFQSSMT

Immunogen

Species (MHC) human (B7)

References De Groot et al. 2001

- The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.
- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes could stimulate IFNγ production in an ELISPOT assay.
- SPAIFQSSMT was confirmed as a previously identified HLA-B7 epitope in this study.

**HXB2 Location** RT (156–165)

**Author Location RT (IIIB)** 

Epitope SPAIFQSSMT

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

Keywords epitope processing, escape

References Moore et al. 2002b

- HIV polymorphisms in the RT protein were examined relation to HLA alleles found in a population of 473 participants in the Western Australian HIV Cohort Study. 64 significant associations between polymorphisms at particular positions and HLA alleles were detected, for HLA-B7, -B12, -B35 and -B15. Fifteen of these were in positions with known epitopes, 4 in anchor residues, 11 in other positions. Six additional polymorphic sites associated with particular HLA molecules flanked known epitopes and may relate to processing.
- HLA-B7+ individuals with a S162x (18/33) substitution had higher viral loads than those that did not, suggesting escape was associated with diminished immune control of viremia.

**HXB2 Location** RT (156–165)

**Author Location** Pol

Epitope SPAIFQSSMT

**Epitope name** 1306

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A03, A24, B07, B38, Cw07, Cw12/13

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for SPAIFQSSMT: 13%

**HXB2 Location** RT (156–165)

**Author Location** RT Pol (311–319)

Epitope SPAIFQSSMT

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/7 patients recognized this epitope.

**HXB2 Location** RT (158-166)

Author Location RT (325-333)

Epitope AIFQSSMTK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

**Keywords** responses in children, mother-to-infant transmission

References Brander & Walker 1995

• Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.

**HXB2 Location** RT (158–166)

**Author Location** RT (325–333 LAI)

Epitope AIFQSSMTK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0301 epitope.

**HXB2 Location** RT (158–166)

**Author Location** 

Epitope AIFQSSMTK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords acute/early infection

**References** Wilson *et al.* 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- $\bullet$  The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.

• No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

HXB2 Location RT (158–166)
Author Location RT (325–333)
Epitope AIFQSSMTK
Immunogen HIV-1 infection
Species (MHC) human (A\*0301)
Keywords immunodominance
References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INFγ responses to other epitopes.
- 1/11 of the A2+ individuals was HLA A3 and reacted with this
  epitope as well as two other A3.1 epitopes.

**HXB2 Location** RT (158–166)

Author Location Pol

**Epitope** AIFQSSMTK **Subtype** A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (A\*0301, A11, A33)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses
  to peptide pools were detected using intracellular cytokine
  staining and IFNgamma Elispot assays after vaccination of 5
  macaques. The response to the Mamu A\*01 SIV p27 epitope
  p11C (CTPYDINQM), included in the polyepitope region, was
  not immunodominant in the Mamu A\*01 vaccinated macaques,
  possibly because of processing limitations in context of the
  artificial polyepitope string Wee et al. [2002].

HXB2 Location RT (158–166)
Author Location RT (325–333 LAI)
Epitope AIFQSSMTK
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (A\*1101)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is an A\*1101 epitope.

HXB2 Location RT (158–166)
Author Location Pol (313–321)
Epitope AIFQSSMTK
Subtype B, CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A\*1101)
Keywords subtype comparisons
References Fukada et al. 2002

- Counterparts for eight known clade B HLA A\*1101 epitopes were generated for clade E (CRF01). Three epitopes, identical among clade A-E, were cross-reactive and recognized by clade E infected individuals. The clade E and B analogs to three more HLA A\*1101 epitopes was recognized in a clade specific manner. Two other HLA A\*1101 clade B defined epitopes were found not to have stimulated a response in clade E infected individuals.
- AIFQSSMTK is commonly found in viruses representing subtypes A-E. It was strongly recognized by CTL from 2/5 B clade infected Japanese subjects, and 5/6 E clade infected Thai subjects.

HXB2 Location RT (158–166)
Author Location RT (313–321)
Epitope AIFQSSMTK
Subtype B, CRF01\_AE
Immunogen
Species (MHC) human (A\*1101)

Country Thailand.

**Keywords** HIV exposed persistently seronegative (HEPS), structure

References Li & Bouvier 2004

• HLA-A\*1101 has been associated with resistance to acquisition of HIV-1 infection in female sex-workers in Thailand. Its crystal structure has been determined in association with two immunodominant A\*1101 HIV-1 CTL epitopes. Its anchor residues are confirmed as P2(Ile/Val) and C-term (Lys). The backbone conformation of the peptides is defined as two bulges separated by a secondary anchor residue (P6 Ser or Met) that may offer various advantages in the selection and presentation of CTL epitopes by HLA-A\*1101.

HXB2 Location RT (158–166) Author Location RT (325–333) Epitope AIFQSSMTK Immunogen HIV-1 infection

**Species (MHC)** human (A\*1101, A3, A\*0301, A\*6801)

**References** Menendez-Arias *et al.* 1998; Threlkeld *et al.* 1997

- Study of the fine specificity of an A3-like super-type epitope (the A3 super-type includes A\*0301, A\*1101, A\*3101, A\*3301, and A\*6801)
- A3 super-type is characterized by a hydrophobic or hydroxyl containing anchor residue at position 2, and a positive charge in the C-term position.

- While most lines were specific, promiscuous cloned CTL lines were also derived from HIV+ donors that could recognize epitope presented by either A3 or A11 or A\*6801.
- · Alanine substitutions throughout the epitope and natural variants indicate that the same amino acid positions are critical for presentation by either MHC molecule, A3 or A11.
- AIFQSSMTK is presented by three members of the A3 superfamily: A\*0301, A\*1101, and A\*6801, and the naturally occurring variants A1S and K9R are recognized with similar efficiency to wild type epitope - AIFQRSMTR can also bind to two additional members of the A3 superfamily, A\*3101 and A\*3301.

**HXB2 Location** RT (158-166)

Author Location RT

Epitope AIFQSSMTK Immunogen HIV-1 infection Species (MHC) human (A11)

References Wagner et al. 1998a

• CTL specific for HIV epitopes were used to show that the mediators of both the cytolytic (granzyme A was used as the marker) and non-cytolytic (HIV-1 inhibitory chemokines MIP-1  $\alpha$  and RANTES were used as markers) anti-viral responses are localized within the CTL's cytotoxic granules.

**HXB2 Location** RT (158-166)

**Author Location** RT (325–333 LAI)

**Epitope** AIFQSSMTK

Subtype B

Immunogen peptide-HLA interaction

Species (MHC) human (A11)

References Menendez-Arias et al. 1998; Zhang et al. • 6/8 tested FSWs recognized this epitope.

• Exploration of A11 binding motif, based on Nixon et al. 1991.

**HXB2 Location** RT (158-166)

**Author Location** RT (325–333 LAI)

Epitope AIFQSSMTK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A11)

Keywords review

References McMichael & Walker 1994

• Review of HIV CTL epitopes.

**HXB2 Location** RT (158–166)

**Author Location** Pol (305–313 93TH253 subtype CRF01)

Epitope AIFQSSMTK Epitope name P313-321 Subtype CRF01\_AE

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A11)

Keywords HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

• This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.

- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers - weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was weakly reactive in the HEPS study subject 128 who was HLA A11/A33.
- This epitope was strongly reactive in HIV+ study subjects 053 and 184 who carried HLA-A11.

**HXB2 Location** RT (158–166)

**Author Location** Pol (305–313 93TH253 subtype CRF01)

**Epitope** AIFQSSMTK Subtype CRF01 AE Immunogen HIV-1 infection Species (MHC) human (A11)

Keywords subtype comparisons

References Bond et al. 2001

- HLA-A11 CRF01 (called subtype E in Bond et al.) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This epitope was predicted by the EpiMatrix method to be likely to bind to A11, and it served as an epitope in the FSWs, it was one of the six A11 epitopes that had been previously defined.
- An HLA-A11 tetramer was made for this epitope, which was recognized by two subjects - and both subjects had expanded tetramer staining T-cell populations after in vitro stimulation.
- This epitope was highly conserved in other subtypes, and exact matches were common.

**HXB2 Location** RT (158–166)

**Author Location** RT (158–166 IIIB)

**Epitope** AIFQSSMTK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

Keywords epitope processing, escape

References Moore et al. 2002b

- HIV polymorphisms in the RT protein were examined relation to HLA alleles found in a population of 473 participants in the Western Australian HIV Cohort Study. 64 significant associations between polymorphisms at particular positions and HLA alleles were detected, for HLA-B7, -B12, -B35 and -B15. Fifteen of these were in positions with known epitopes, 4 in anchor residues, 11 in other positions. Six additional polymorphic sites associated with particular HLA molecules flanked known epitopes and may relate to processing.
- HLA-A11+ individuals with a K166x (4/19) substitution had higher viral loads than those that did not, suggesting escape was associated with diminished immune control of viremia.

**HXB2 Location** RT (158–166)

**Author Location** Pol

Epitope SIFQSSMTK
Immunogen HIV-1 infection
Species (MHC) human (A11)

**Donor MHC** A2, A11, B8, B60, Bw6

**Keywords** HAART, ART **References** Appay *et al.* 2002

- Four HIV patients with prolonged clinically successful antiviral therapy but with ongoing evidence of replication and Nef mRNA transcription, showed specific T-cell responses by Elispot and Tetramer staining, maintained for 2-4 years after initiation of HAART.
- Nef epitope recognition was detected in all 4 subjects, gp120,
   Pol and Gag-specific in 1 or 2 subjects.
- The HIV-specific CD8+ T-cells had an intermediate maturation phenotype characterized by low levels of perforin and high levels of CD27 expression.

**HXB2 Location** RT (158–166)

Author Location Pol (314-322)

Epitope AIFQSSMTK

Epitope name AK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

**Keywords** optimal epitope **References** Allen *et al.* 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For one of the escape variants, a novel CD8 T-cell response equal in magnitude to the wild type, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wild type.
- This epitope did not vary.

**HXB2 Location** RT (158–166)

Author Location Pol (314–322)

Epitope AIFQSSMTK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A11)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** RT (158–166)

Author Location Pol (313-321)

**Epitope** AIFQSSMTK

**Epitope name** AK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

**Donor MHC** A\*01, A\*11, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords escape, optimal epitope

References Koibuchi et al. 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- The AK9 variant AIFQSSMTr was essentially the only form of the epitope detected over a 5-year period in this person. Elispot reactions indicated the T-cell clones only recognized the autologous form, not the B clade consensus, AIFQSSMTK. Two rare variants were observed at the 5-year time point, tIFQSSMTr and AIFQSSMar.

**HXB2 Location** RT (158–166)

Author Location Pol (325–333)

Epitope AIFQSSMTK

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (A11, A\*0301, A33)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot

granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- One of the patients responded to this peptide with GzB producing cells, while none of the patients responded with IFN-gamma producing cells.

**HXB2 Location** RT (158–166)

Author Location RT (B consensus)

**Epitope** AIFQSSMTK

Epitope name ATK9

Immunogen HIV-1 infection Species (MHC) human (A11, A3)

**Donor MHC** A02, A11, B18, B44, Cw5, Cw12; A03, B14,

B60, Cw3, Cw7; A01, A03, B08, B14, Cw7,

Cw8

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, crosspresentation by different HLA, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
   activation in dying target cells, it was shown that the subset of
   HIV-1-specific CD8+ T cells secreting both IFN-gamma and
   TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1 specific CD8+ T-cell maturation phenotypes and intracellular
   perforin expression was found.
- 3/9 individuals recognized this epitope, two presented by HLA-A3, one presented by HLA-A11.

**HXB2 Location** RT (158–166)

Author Location RT (325–333 IIIB)

**Epitope** AIFQSSMTK **Immunogen** HIV-1 infection

Species (MHC) human (A3)

**Keywords** responses in children, mother-to-infant transmission

References Wilson et al. 1996

- Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.
- AIFQSSMTR and AILQSSMTK, naturally occurring variants, were found in infant, and are recognized.
- TISQSSMTK, a naturally occurring variant, was found in infant and is not recognized.

**HXB2 Location** RT (158–166)

**Author Location** RT (325–333 LAI)

Epitope AIFQSSMTK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Keywords** subtype comparisons **References** Cao *et al.* 1997a

- The consensus peptide of B and D clade viruses is 2 to 17 epitopes were recognized in a given individual, A2-AIFOSSMTK.
- The consensus peptide of a subset of As is AIFQASMTK and it is less able to stimulate the CTL clone.
- The consensus peptide of a subset of As is SIFQSSMTK and is as reactive as the originally defined epitope.

**HXB2 Location** RT (158–166)

**Author Location** Pol (325–333 IIIB)

Epitope AIFQSSMTK
Immunogen HIV-1 infection
Species (MHC) human (A3)

**Keywords** responses in children, mother-to-infant transmission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- One variant found in an infant gave a positive CTL response: AIFQSSMTR.

• AIFLSSMTK and TISQSSMTK were escape mutants.

**HXB2 Location** RT (158–166)

**Author Location** RT (325–333 SF2)

**Epitope** AIFQSSMTK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A3+ individuals that had a CTL response to this epitope broken down by group: 0/7 group 1, 0/4 group 2, and 1/2 group 3.

**HXB2 Location** RT (158–166)

Author Location RT (158-166)

Epitope AIFQSSMTK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- All patients recognized at least 1 A3 epitope, up to 8 A3 epitopes, but none was clearly dominant.
- In two of the subjects, AIFQSSMTK was the dominant epitope.

**HXB2 Location** RT (158–166)

Author Location RT Pol (313–321)

Epitope AIFQSSMTK

**Epitope name** A3-ATK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

 CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.

- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 3/7 individuals began to have detectable responses to this epitope after STI.

**HXB2 Location** RT (158–166) **Author Location** RT (158–166)

**Epitope** AIFQSSMTK **Epitope name** A3-AK9 Pol

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

- An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.
- The second infecting strain had the variant aifqssmlk. The CTL response to the second variant was zero or low at all timepoints.
   The CTL response to the first variant was also low, and declined over time.

**HXB2 Location** RT (158–166)

**Author Location** RT (158–166)

Epitope AIFQSSMTK
Immunogen HIV-1 infection
Species (MHC) human (A3)

**Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4, DR8, DO7

DKo, DQ/

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these epitopes, but this increased over time.

HXB2 Location RT (158–166) Author Location RT Pol (313–333) Epitope AIFQSSMTK Immunogen HIV-1 infection Species (MHC) human (A3)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/14 patients recognized this epitope.

**HXB2 Location** RT (158-166)

**Author Location** Pol

Epitope AIFQSSMTK

**Epitope name** AK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 9, aifqssmtI, was found not to correspond to the most polymorphic residues in the epitope.

**HXB2 Location** RT (158–166)

**Author Location** RT (325–333 LAI)

Epitope AIFQSSMTK

Epitope name P3

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (A3 supertype)

Keywords HAART, ART, supertype

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+PBL but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

**HXB2 Location** RT (158–166) Author Location Pol (337–345) **Epitope** AIFQSSMTK Immunogen HIV-1 infection **Species (MHC)** human (A3 supertype) Keywords supertype, rate of progression References Propato et al. 2001

• Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer

epitopes.

• Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.

- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, Among HLA-A3 women, 2/2 HEPS and 3/3 HIV-1 infected A\*1101, A\*3101, A\*3301 and A\*6801).

**HXB2 Location** RT (158-166)

**Author Location** Pol

Epitope AIFQSSMTK

Epitope name 1339 Subtype multiple Immunogen HIV-1 infection

**Species (MHC)** human (A3, A\*0301, A11, A\*6801, A33)

Donor MHC A02, A03, B08, B51, Cw01, Cw07; A03, A26, B08, B52; A03, A11, B14, B05, Cw08

Country United States. Assay type T-cell Elispot

Keywords binding affinity, supertype, computational epitope prediction, cross-presentation by differ-

ent HLA

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for AIFQSSMTK: 59% Supertype epitope binding to A03, A3.1, A11, A6801, A33.

**HXB2 Location** RT (158–166) Author Location Pol (313-321) **Epitope** AIFQSSMTK Immunogen HIV-1 infection Species (MHC) human (A3, A11) References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** RT (158–166) Author Location Pol (325–333) **Epitope** AIFQSSMTK

**Immunogen** HIV-1 infection, HIV-1 exposed seronegative • HIV-1 specific CTLs release  $\gamma$ -IFN, and  $\alpha$ - and  $\beta$ -TNF.

Species (MHC) human (A3, A11, A33)

Keywords HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- Variants (S/A)IFQSSMTK are specific for the A/B clades.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in one of the 2/2 HEPS cases and in one of the 3/3 HIV-1 infected women.

**HXB2 Location** RT (158–166)

**Author Location** RT (325–333 LAI)

**Epitope** AIFQSSMTK

Subtype B

Immunogen

Species (MHC) human (A33)

References Rowland-Jones 1995

· Defined as minimal peptide by titration curve, S. Rowland-Jones, pers. comm.

**HXB2 Location** RT (158–166)

**Author Location** 

Epitope AIFQSSMTK Immunogen HIV-1 infection Species (MHC) human (A33)

> Keywords HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted - 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls, ML1668.

**HXB2 Location** RT (158–182)

**Author Location** RT (325–349 PV22)

Epitope AIFQSSMTKILEPFRKQNPDIVIYQ

Immunogen HIV-1 infection Species (MHC) human (A11)

References Jassoy et al. 1993

HXB2 Location RT (158–182) Author Location RT (325–349)

Epitope AIFQSSMTKILEPFRKQNPDIVIYQ

Immunogen HIV-1 infection Species (MHC) human (A11) References Price *et al.* 1995

• Study of cytokines released by HIV-1 specific activated CTL.

HXB2 Location RT (164–172)
Author Location Pol (343–351)
Epitope MTKILEPFR
Immunogen HIV-1 infection
Species (MHC) human (A3 supertype)
Keywords supertype, rate of progression
References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 4/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location RT (173–181) Author Location RT (173–181 LAI) Epitope KQNPDIVIY Subtype B

Immunogen

Species (MHC) human (A\*3002)

Keywords optimal epitope

References Frahm et al. 2007; Goulder et al. 2001a

• C. Brander notes this is an A\*3002 epitope.

**HXB2 Location** RT (173–181)

**Author Location RT** 

Epitope KQNPDIVIY
Epitope name KY9 (RT-53)
Immunogen HIV-1 infection
Species (MHC) human (A\*3002)

References Goulder et al. 2001a

- HLA-A\*3002 is very common in African populations, 50% of Zimbabweans express HLA-A30, 44% in African Zulu, so five new HIV epitopes were characterized that are presented by this HLA molecule.
- A rapid method was developed combining ELISPOT with intracellular IFN-γ staining of PBMCs to map optimal epitopes, then HLA presenting molecules were defined this method was completed within 48 to 72 hours of receipt of blood.
- Two individuals were studied: Subject 199 (HLA A\*0201/\*3002 B\*4402/51 Cw2/5), a Caucasian, and Subject 6007 (HLA A\*3002/ B53/\*5801 Cw4/7) an African-Caribbean.
- In both HLA-A\*3002 individuals the response to RSLYNT-VATLY was dominant.

- In subject 199 four additional A\*3002 epitopes were identified.
- Three quantitative assays, ELISPOT, precursor frequency and chromium release, confirmed a hierarchy of response: RY11 (p17) > KY9 (gp41) > KY9 (RT-53) > IY9 (gp41)

**HXB2 Location** RT (173–181)

**Author Location** (C consensus)

Epitope AQNPEIVIY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Country South Africa.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** RT (173–183)

**Author Location** (C consensus)

Epitope AQNPEIVIYQY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- AONPEIVIYOY is an optimal epitope.

**HXB2 Location** RT (175–183)

Author Location (C consensus)

Epitope NPEIVIYQY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*18)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

sociated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** RT (175–183) **Author Location** (C consensus) Epitope NPEIVIYQY Subtype C Immunogen HIV-1 infection Species (MHC) human (B\*35) Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- NPEIVIYQY is an optimal epitope.

**HXB2 Location** RT (175–183) **Author Location** RT (328–336 IIIB) **Epitope NPDIVIYQY** Immunogen HIV-1 infection Species (MHC) human (B\*3501) References Tomiyama et al. 1997

- A CTL clone responsive to this epitope was obtained.
- 3/7 B35-positive individuals had a CTL response to this epi-
- D to E, or V to I, substitutions at positions 3 or 5, respectively, reduces CTL activity and binding to B\*3501.

**HXB2 Location** RT (175–183) **Author Location** RT (328–336 IIIB) **Epitope NPDIVIYQY** Immunogen HIV-1 infection Species (MHC) human (B\*3501) Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is a B\*3501 epitope.

**HXB2 Location** RT (175–183) **Author Location** RT (342–350 LAI) Epitope HPDIVIYQY Subtype B Immunogen HIV-1 infection

Species (MHC) human (B\*3501) Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is a B\*3501 epitope.

**HXB2 Location** RT (175–183) Author Location Pol (330–338) Epitope NPDIVIYQY Immunogen HIV-1 infection Species (MHC) human (B\*3501) References Tomiyama et al. 2000a

• CD8+ T-cells that bound one of six HIV-specific B\*3501epitope tetramers did not express CD28 or CD45A.

- Mutational patterns in the I4 residue of NPEIVIYOY are asof CD28+CD45RA+ cells was observed in chronically HIV-1infected individuals relative to healthy individuals.
  - CD28-CD45RA- cells are likely to be effector cells and have high levels of perforin in their cytoplasm.
  - The mean percentage of total CD28- CD8+ cells in chronically infected HIV-1-infected patients was 76.6% in comparison to HIV-1-uninfected individuals (40.6%)

**HXB2 Location** RT (175–183) **Author Location** RT (175–183 IIIB) Epitope NPDIVIYQY Subtype B Immunogen HIV-1 infection

Species (MHC) human (B\*3501) Keywords epitope processing, escape

References Moore et al. 2002b

- HIV polymorphisms in the RT protein were examined relation to HLA alleles found in a population of 473 participants in the Western Australian HIV Cohort Study. 64 significant associations between polymorphisms at particular positions and HLA alleles were detected, for HLA-B7, -B12, -B35 and -B15. Fifteen of these were in positions with known epitopes, 4 in anchor residues, 11 in other positions. Six additional polymorphic sites associated with particular HLA molecules flanked known epitopes and may relate to processing.
- NPDIVIYOY was one of two epitopes characterized in detail. D177x substitutions are know to specifically abrogate binding to HLA-B\*3501, and not other B\*35 subtypes. D177x substitutions were associated with people who carried HLA-B\*3501 and not other B\*35 subtypes; considering high resolution typing generally strengthened the B\*35 associations.

**HXB2 Location** RT (175–183) **Author Location** RT (175–183) Epitope NPDIVIYQY Subtype B Immunogen HIV-1 infection Species (MHC) human (B\*3501) Donor MHC A\*2301, B\*3501, B\*1503, Cw2, Cw7

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, earlyexpressed proteins

References Cao et al. 2003

• All proteins were scanned and optimal epitopes were mapped in a study of CD8+ INFy T-cell responses in 21 men within 15-92days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN $\gamma$ secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.

- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized: 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

**HXB2 Location** RT (175–183) Author Location Pol (330–338)

Epitope HPDIVIYQY

Epitope name HY9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*3501)

**Donor MHC** A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNy, Intracellular cytokine staining, Chromium-release assay

Keywords escape, acute/early infection

References Bansal et al. 2005

- · Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- The response to this epitope was not detected until month 25.

**HXB2 Location** RT (175–183)

**Author Location** 

Epitope NPEIVIYQY

Epitope name NY9

Immunogen

Species (MHC) human (B18)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B18 epitope.

**HXB2 Location** RT (175–183)

**Author Location** RT (342–350 LAI)

Epitope HPDIVIYQY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

Kevwords review

References McMichael & Walker 1994

• Review of HIV CTL epitopes.

**HXB2 Location** RT (175–183)

**Author Location** RT (329–337)

Epitope HPDIVIYQY

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Rowland-Jones et al. 1995

NPDVILIQY is also recognized.

**HXB2 Location** RT (175–183)

**Author Location (SF2)** 

Epitope NPDIVIYQY

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords binding affinity, rate of progression, escape References Kawana et al. 1999

- HLA B35 is associated with rapid disease progression.
- The sequences of 9 previously described HIV-1 B35 CTL epitopes were obtained in 10 HLA B35+ and 19 HLA B35- individuals.
- 3/9 CTL epitopes had substitutions that were more common in B35+ individuals than in B35- individuals – only one of these reduced the binding of the peptide to B35 and was shown to be an escape mutation.
- npEiviyqy was found in 8/10 of the B35+ individuals, and two of the B35- individuals—the D->E substituted peptide had reduced binding affinity to B35 and may be an escape mutant.

**HXB2 Location** RT (175–183)

Author Location RT (329–337)

Epitope HPDIVIYQY

Immunogen in vitro stimulation or selection

Species (MHC) human (B35)

References Lalvani et al. 1997

- A peptide-based protocol was optimized for restimulation of CTLp using optimized peptide and IL-7 concentrations – importantly this protocol does not stimulate a primary response, only secondary - peptide-specific CTLp counts could be obtained via staining with peptide-Class I tetramers.
- This peptide was one of the B35 presented test peptides used in control experiments showing that the assay gave no activity using lymphocytes from 21 healthy B35 seronegative donors.

**HXB2 Location** RT (175-183)

**Author Location** RT (328–336 IIIB)

Epitope NPDIVIYQY

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Menendez-Arias et al. 1998; Shiga et al. 1996

- Binds HLA-B\*3501.
- · CTL activity to this epitope was originally detected in a longterm survivor, however it has since been found in normal progressors - it is cross-reactive with HIV-2 (HPDILIYQY), but D3E and V5I substitutions reduce binding Menendez-Arias et al. [1998]

**HXB2 Location** RT (175–183)

**Author Location** RT (328–336 IIIB)

Epitope NPDIVIYQY

**Immunogen** HIV-1 infection

Species (MHC) human (B35)

Keywords review, escape

References Menendez-Arias et al. 1998; Sipsas et al.

1997

- HIV IIIB proteins were used to define the range of CTL epitopes recognized by 3 lab workers accidentally infected with HIV-1 IIIB.
- NPDIIIYQY, a variant found in HIV-1 JRCSF, was also recognized.
- NPEIVIYQY, was also recognized.
- · NPDLVIYQY, was also recognized.
- Menendez-Arias et al. [1998], in a review, notes that the YXDD motif, highly conserved among polymerases, overlaps this epitope CTL activity to this epitope was originally detected in a long-term survivor, however it has since been found in normal progressors it is cross-reactive with HIV-2 (HPDILIYQY), but D3E and V5I substitutions reduce binding.

**HXB2 Location** RT (175–183)

**Author Location RT** 

Epitope NPDIVIYQY

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B35)

**Keywords** review, subtype comparisons, HIV exposed persistently seronegative (HEPS)

**References** Menendez-Arias *et al.* 1998; Rowland-Jones *et al.* 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades such cross-reactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A subtype consensus is HPDIVIYQY.
- The D subtype consensus is NPEIVIYQY.
- Menendez-Arias et al. [1998], in a review, notes that the YXDD motif, highly conserved among polymerases, overlaps this epitope CTL activity to this epitope was originally detected in a long-term survivor, however it has since been found in normal progressors it is cross-reactive with HIV-2 (HPDILIYQY), but D3E and V5I substitutions reduce binding.

**HXB2 Location** RT (175–183)

Author Location Pol (subtype B)

Epitope NPDIVIYQY

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B35)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- Clade A version of epitope HPDIVIYQY, Clade D NPEIVIYQY.

**HXB2 Location** RT (175–183)

Author Location Pol

Epitope HPDIVIYQY

Immunogen

Species (MHC) human (B35)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied – these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive, and the B35 allele seems to be protective.
- HIV-2 version of this epitope is not conserved: NPDVILIQY, but the CTLs are cross-reactive – one of five B35 CTL epitopes that are cross-reactive, see also Rowland-Jones et al. [1995]

**HXB2 Location** RT (175-183)

**Author Location** 

Epitope HPDIVIYQY

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords acute/early infection

References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

**HXB2 Location** RT (175–183)

**Author Location** Pol (subtype A)

Epitope HPDIVIYQY

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (B35)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- HPDIVIYQY or NPDIVIYQY was recognized in 1 of the 6 women (ML857), and the response was present in the last available sample prior to seroconversion, 7 months.

- 20/20 sequences of the infecting strain had three substitutions in this epitope, all 20 were NpQiIiyqy, and this form was not recognized by CTL from ML 857 this was the only case in the study where a virus carrying an unrecognized form of the epitope broke through.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- NPDIVIYQY was recognized by 1/22 HEPS control sex workers, ML887.

**HXB2 Location** RT (175–183)

Author Location RT (175–183 SF2)

Epitope NPDIVIYQY Immunogen HIV-1 infection Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B35+ individuals that had a CTL response to this epitope broken down by group: 1/2 group 1, 0/2 group 2, and 1/1 group 3.

**HXB2 Location** RT (175–183)

**Author Location** Pol (342–350)

Epitope HPDIVIYQY

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B35)

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

- Variants (H/N)PDIVIYOY are specific for the A/B clades.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B35 women, 2/3 HEPS and 1/4 HIV-1 infected women recognized this epitope.

- The dominant response to this HLA allele was to this epitope in only one of the 2/3 HEPS cases, and was not to this epitope in the one responsive HIV-1 infected women.
- Subject ML 857 shifted from a A\*6802 DTVLEDINL and B35 (H/N)PDIVIYQY response prior to seroconversion to a B35 PPIPVGDIY and B35 VPLRPMTY response postseroconversion and the loss of the pre-seroconversion response was not due to sequence variation within these epitopes.

**HXB2 Location** RT (175-183)

**Author Location** 

Epitope HPDIVIYQY

Epitope name Pol-HY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA B35, 4/21 (19%) recognized this epitope.

**HXB2 Location** RT (175–183)

**Author Location** Pol

Epitope NPDIVIYQY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

**Donor MHC** A3, A11, B35, B51

Keywords mother-to-infant transmission

References Sabbaj et al. 2002

- IFNgamma T-cell responses in breast milk of 5 HIV-1 infected women from the US and 6 from Zambia were tested with using Elispot. 11/11 women responded to Gag, 8/11 responded to Pol, 7/11 women to Nef, and 2/5 women to Env peptide pools. These responses were shown to be primarily due to CD8+ T-cells in one woman, and another woman had cytolytic responses measured by Cr-release.
- T-cells in breast milk from a volunteer who was HLA A3, A11, B35, B51 induced IFNgamma after stimulation with a peptide that carries known B35 epitope NPDIVIYQY.
- The frequencies of responses in the two compartments differed, and 2/4 women that responded to epitopes in Nef 101-205 and Pol 601-710 showed responses in breast milk but no detectable responses in peripheral blood cells.

**HXB2 Location** RT (175–183)

**Author Location** Pol

Epitope HPDIVIYQY

Subtype A

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade

HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (B35)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

**HXB2 Location** RT (175-183) Author Location Pol (342–350) Epitope HPDIVIYQY Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35) Country United States.

> Assay type CD8 T-cell Elispot - IFNγ, CD8 T-cell Elispot granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- None of three patients responded to this peptide with GzB producing cells, while one of the patients responded with IFNgamma producing cells.

**HXB2 Location** RT (175–183) **Author Location** RT Pol (330–338) Epitope HPDIVIYQY Immunogen HIV-1 infection Species (MHC) human (B35)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/9 patients recognized this epitope.

**HXB2 Location** RT (175-184)

**Author Location** RT (175–184 LAI) Epitope NPDIVIYQYM Subtype B Immunogen HIV-1 infection

Species (MHC) human (B51) References Samri et al. 2000 This epitope contains the mutation M184V, a frequent mutation

- induced by nucleoside reverse transcriptase inhibitors. Patient 246#1 (B51), was found by ELISPOT to recognize the
- wild type and the mutated peptide after zidovudine treatment. • The resistance mutation M184V gave binding increased predicted score to B51 (http://bimas.dcrt.nih.gov/molbio/hla\_bind) compared to the wildtype RT sequence and also an increased ELISPOT

**HXB2 Location** RT (175–199) **Author Location** RT (342–366 LAI) Epitope NPDIVIYQYMDDLYVGSDLEIGQHR

Subtype B

reactivity.

Immunogen HIV-1 infection Species (MHC) human (A11)

References Menendez-Arias et al. 1998; Walker et al. 1989

• One of five epitopes defined for RT-specific CTL clones in this study.

**HXB2 Location** RT (179-187)

**Author Location RT** 

Epitope VIYQYMDDL

Immunogen vaccine

Vector/Type: vaccinia

Species (MHC) human (A\*0201)

References Hanke et al. 1998a; Hanke et al. 1998b

• This epitope was shown to be processed and presented to appropriate CTL clones upon infection of human target cells with vaccinia virus Ankara (VVA) carrying 20 HIV-1 epitopes recognized by humans.

**HXB2 Location** RT (179–187)

**Author Location RT** 

Epitope VIYQYMDDL

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Tan et al. 1999

- Adoptive transfer of two autologous in vitro-expanded CTL clones against the A\*0201 restricted epitopes SLYNTVATL and VIYQYMDDL were infused into a patient - they were well tolerated, but the SLYNTVATL clone was shown by tetramer staining to be rapidly eliminated through apoptosis, and the treatment had no impact upon viral load and CD4 and CD8 cell
- Tetramer staining failed for the VIYQYMDDL epitope as the tetramer was unstable.

**HXB2 Location** RT (179–187) Author Location Pol (346-354) Epitope VIYQYMDDL **Immunogen** HIV-1 infection

Species (MHC) human (A\*0201)

cape

References Sewell et al. 1999

- Proteasome regulation influences epitope processing and could influence patterns of immunodominance.
- The proteasome is inhibited by lactacystin treatment, and gamma IFN induces expression of proteasome subunits, LMP2 and LMP7, which combine with the proteasome to create an immunoproteasome.
- IFN-gamma induction of the immunoproteasome and lactacystin inhibition increases the presentation of the A\*0201 VIYQYMDDL epitope, but decreases the presentation of the A\*0201 ILKEPVHGV epitope, which is immunodominant within pol proteins, showing the two epitopes are processed by different pathways.
- ILKEPVHGV seems to be processed by the classical proteasome pathway, while VIYQYMDDL appears to be destroyed by this pathway.
- This epitope contains the catalytic site (YMDD) of RT, a conserved sequence in HIV-1 which restricts escape mutants.

**HXB2 Location** RT (179–187)

**Author Location** RT (346–354 LAI)

Epitope VIYQYMDDL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Kevwords review

References Harrer et al. 1996a; Menendez-Arias et al.

- The substitution VIYQYVDDL abrogates CTL response and confers drug resistance.
- Menendez-Arias et al. [1998], in a review, notes that this epitope includes catalytic residues (Asp-185 and Asp-186) in the active site of RT.

**HXB2 Location** RT (179–187)

**Author Location** RT (346–354 LAI)

Epitope VIYQYMDDL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0201 epitope.

**HXB2 Location** RT (179–187)

Author Location RT (346-354)

Epitope VIYQYMDDL

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords review, escape

References Brander et al. 1998a; Menendez-Arias et al.

- Of 17 infected HLA A\*0201 subjects, 13 had CTL responses against the p17 SLYNTVATL epitope, six recognized ILKEPVHGV and five recognized VIYQYMDDL, and there was no correlation between viral load and recognition of a specific epitope or evidence of immune escape.
- · Only one subject had CTL against all three epitopes.

- **Keywords** epitope processing, immunodominance, es• Subjects were part of the San Francisco City Clinic Cohort, the ARIEL project and from the Boston area.
  - In the review Menendez-Arias et al. [1998] the authors note that substitution of three residues in this epitope can confer resistance to RT inhibitors (1, 3, and 6) – substitutions V1E and M6V abolish CTL activity, and M6V confers resistance to 3TC - substitution Y3C reduces CTL activity and is associated with resistance to non-nucleoside RT inhibitors.

**HXB2 Location** RT (179–187)

**Author Location RT** 

Epitope VIYQYMDDL

Epitope name RT VL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords subtype comparisons, supertype, computational epitope prediction

References Altfeld et al. 2001c

- HIV was scanned for all peptides which carried the A2supermotif pattern conserved in more than 50% of B clade sequences – 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 - 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.
- Three additional previously described HLA-A2 epitopes were added to the set of 20, including RT VL9, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- RT VL9 was not recognized by any of the 22 HLA-A2 patients with chronic HIV-1 infection or the 13 HLA-A2 patients with acute HIV-1 infection included in this study.

**HXB2 Location** RT (179-187)

**Author Location** RT (346–354)

Epitope VIYQYMDDL

Epitope name VL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Dela Cruz et al. 2000

- Integration of HIV RT CTL epitopes into the N-terminus of the HLA-A2 heavy chain, or tethering the epitopes to the target chain, resulted in epitope-specific lysis by CD8+ CTL.
- These antigens could also be used to stimulate primary responses in vitro.

**HXB2 Location** RT (179-187)

**Author Location** Pol (346–354)

Epitope VIYQYMDDL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords epitope processing, immunodominance

References Sewell et al. 2002

• Epitope processing of three different HLA-A\*0201 HIV epitopes was shown to use different pathways, which might influence patterns of immunodominance. .174 cells were used that lack TAP1 and TAP2 genes, as well as the LMP2 and LMP7 genes that encode the beta-subunits of the immunoproteasome. These genes could be added back through transfection to study processing.

ILKEPVHGV was efficiently presented in TAP-1 and -2 transfected cells while VIYQYMDDL and SLYNTVATL were not. VIYQYMDDL was destroyed by the MB1 subunit of the protease, and could be expressed in the presence of the proteasome inhibitor lactacystin, but SLYNTVATL expression was not restored. SLYNTVATL expression was unaltered by lactacystin in a wild type cell line.

**HXB2 Location** RT (179–187)

**Author Location** RT (346–354 LAI)

Epitope VIYQYMDDL

Epitope name LR26

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant (IFA), Montanide (ISA 720), P30, PLG

Species (MHC) mouse (A\*0201)

Keywords binding affinity, vaccine-specific epitope char-

acteristics, immunodominance

References Peter et al. 2001

- The stability of peptide binding to HLA-A2.1 was determined for six HLA-A2.1 peptides included in this vaccine study

   ILKEPVHGV (RT), SLYNTVATL (p17), SLLNATDIAV (gp41) and LLWKGEGAV (RT) all bound with high affinity comparable to a influenza epitope reference (GILGFVFTL), while RGPGRAFVTI and VIYQYMDDL bound with a lower affinity (relative binding activity = 0.01).
- The four high-affinity peptides formed stable complexes with half-lives ranging between 8 and 32 hours, while the low affinity peptides had half lives of less than an hour.
- HLA-A2.1 transgenic mice were immunized with the six HIV-1 peptides and P30, as a universal T-helper epitope, with IFA or Montanide or microspheres as adjuvants.
- All peptides except VIYQYMDDL induced a stong CTL response in Cr-release assays stronger responses were observed when peptides were delivered alone, indicating immunodominance when the combination was used.

**HXB2 Location** RT (179–187)

**Author Location** RT (346–354 LAI)

Epitope VIYQYMDDL

Epitope name LR26

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant

(IFA), IL-12, P30

Species (MHC) mouse (A\*0201)

Keywords vaccine-specific epitope characteristics, im-

munodominance

References Peter et al. 2002

When HIV-1 peptides were used to vaccinate HLA-A2.1 transgenic A2-Kb mice, strong responses to five peptides were observed when the peptides were given individually, but immunodominance limited the response to some of the peptides when they were given in combination Peter et al. [2001]. IL-12 can counteract immunodominance in BALB/c mice, so it was given

with the multiple epitope vaccination, and was instead found to specifically eliminate the HLA-A2.1-epitope CTL responses, but not Kb CTL responses. This was possibly a consequence of transient depletion of T-cells, B cells and macropahges in the spleen.

**HXB2 Location** RT (179-187)

**Author Location** Pol

**Epitope** VIYQYMDDL **Subtype** A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

**Species (MHC)** human, macaque (A\*0201)

Keywords subtype comparisons, epitope processing,

vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the A-clade epidemic in Nairobi, Kenya. A DNA and MVA prime-boost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

**HXB2 Location** RT (179–187)

Author Location RT (179-187)

Epitope VIYQYMDDL

Subtype B

Immunogen vaccine

Vector/Type: peptide HIV component: RT Adjuvant: Incomplete Freund's Adjuvant

(IFA), IL-12

Species (MHC) mouse (A\*0201)

Donor MHC A2.1

Assay type Cytokine production, Chromium-release as-

Keywords binding affinity, vaccine-induced epitopes

References Okazaki et al. 2003

Alanine substitutions of VIYQYMDDL were tested for importance of each amino acid for HLA-A2.1 binding. Peptide variant (vLyqymddV) showed an 8 fold higher MHC binding affinity than wild type. YLyqymddV had an even higher binding affinity, but the Y at position one blocked TCR recognition. The higher affinity form of vLyqymddV induced CTL in vivo

that could protect against a vaccinia virus expressing RT and the wild type epitope.

 $\begin{array}{ll} \textbf{HXB2 Location} & RT~(179\text{--}187) \\ \textbf{Author Location} & RT~(179\text{--}187~MN) \end{array}$ 

Epitope VIYQYMDDL

Subtype B

Immunogen vaccine

Vector/Type: DNA, polyepitope Strain: B clade MN HIV component: gp120, Protease, RT Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) humanized mouse (A\*0201) Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance, immunotherapy

References Isaguliants et al. 2004

- Immunization of HLA-A\*0201-transgenic mice with synthetic genes encoding clusters of human A\*0201 CTL epitopes located at the sites of drug resistance mutations, induced RTspecific cellular responses indicating the immunogenicity of these constructs. This vaccine strategy may be a first step towards a therapeutic vaccine against drug-resistant strains.
- This was one of five HLA-A\*0201 epitopes from the RT or protease proteins that was included in the polyepitope vaccine. When the transgenic HLA A\*0202 mice were vaccinated with the polyepitope construct or with a mixture of RT peptides, a sustained low level CD8+ T-cell gamma IFN response was observed, in contrast to when an intact RT gene was used for vaccination.

 $\textbf{HXB2 Location} \ \ RT\ (179\text{--}187)$ 

**Author Location** Pol (346–354)

Epitope VIYQYMDDL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- One of seven patients responded to this peptide with GzB producing cells, while none of the patients responded with IFN-gamma producing cells.

**HXB2 Location** RT (179–187)

**Author Location** (C consensus)

Epitope VIYQYMDDL

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, optimal epitope **References** Kiepiela *et al.* 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- VIYQYMDDL is an optimal epitope.

**HXB2 Location** RT (179–187)

**Author Location RT** 

Epitope VIYQYMMDL

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2)

**Keywords** subtype comparisons, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades such cross-reactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A and D consensus sequences are both VIYQYMMDL.

 $\textbf{HXB2 Location} \ \ RT \ (179\text{--}187)$ 

Author Location Pol (346-354)

Epitope VIYQYMDDL

Immunogen vaccine

Vector/Type: DNA prime with vaccinia boost

Species (MHC) human (A2)

References Woodberry et al. 1999

- A polyepitope vaccine was generated in a vaccinia construct that contiguously encoded seven epitopes, all presented by HI A A-2
- HHD mice have a transgene of HLA A2 linked to the transmembrane and cytotoxic domains of H-2D<sup>d</sup> – this transgene is the only MHC molecule expressed in the mice.
- CTL responses to Gag (77-85) SLYNTVATL, Pol (476-484) ILKEPVHGV, gp120 (120-128) KLTPLCVTL, and Nef (190-198) AFHHVAREL were observed in HIV polytope HHD-vaccinated mice, and these responses were enhanced with vaccinia boost.
- No CTL immune responses were generated against HLA A2-restricted HIV epitopes Nef 157-166 (PLTFGWCYKL), Pol 346-354 (VIYQYMDDL), and Nef 180-189 (VLEWRFDSRL)
- Sixteen HLA A2+ patients were tested for their ability to make CTL responses by peptide restimulation in culture with the epitopes selected for inclusion in the polytope one individual recognized all seven of these epitopes; 7 patients had CTL cultures able to recognize at least one of the epitopes, and 6 of those 7 recognized more than one epitope, but they were not able to test all peptides for all patients; many patients only had three peptides tested.
- VIYQYMDDL was recognized by 3 of the HLA-A2 patients.

**HXB2 Location** RT (179–187)

**Author Location** RT (179–187)

Epitope VIYQYMDDL

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** escape, immunotherapy References Schmitt et al. 2000

- The mutation M184V confers resistance to lamivudine, and is in the middle of the HLA-A2 epitope VIYQYMDDL.
- 1/28 individuals tested produced HIV-1 RT-specific CTL that recognized the peptide representing the lamivudine escape mutants VIYQYVDDL and VIYQYIDDL, but failed to recognize the wildtype epitope VIYOYMDDL.
- This suggests immunotherapy stimulating anti-VIYQYVDDL responses maybe helpful for reducing lamivudine escape.

**HXB2 Location** RT (179–187) **Author Location** RT (179–187) Epitope VIYQYMDDL Immunogen HIV-1 infection Species (MHC) human (A2) References Haas et al. 1998

• Of 98 patients in cross-sectional analysis, 78% had CTL against pol – RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)

**HXB2 Location** RT (179–187)

**Author Location** Pol (339–347 93TH253 subtype CRF01)

**Epitope** VIYQYMDDL Epitope name P334-342 Subtype CRF01 AE Immunogen HIV-1 infection Species (MHC) human (A2)

> Keywords HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers - weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not
- This epitope was reactive in HIV+ control study subject 144 who carried HLA-A2.

**HXB2 Location** RT (179–187)

**Author Location** Pol (339–347 93TH253 subtype CRF01)

Epitope VIYQYMDDL Subtype CRF01\_AE Immunogen HIV-1 infection Species (MHC) human (A2) **Keywords** subtype comparisons References Bond et al. 2001

• More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes

were also tested.

- 2/4 tested FSWs recognized the E clade version of this epitope, which is identical to the previously defined B clade version VIYOYMDDL.
- This epitope was conserved in many subtypes, and exact matches were very uncommon.

**HXB2 Location** RT (179–187) Author Location RT (179-187)

Epitope VIYQYMDDL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords rate of progression, acute/early infection References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.

**HXB2 Location** RT (179-187)

Author Location Pol (346–354 LAI)

**Epitope** VIYQYMDDL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords HAART, ART, epitope processing

References Kelleher et al. 2001a

- Ritonavir (RTV) inhibits chymotryptic activity in the 20S proteasome in vitro, as does Saquinavir (SQV) to a lesser extent; Indinavir (IDV) does not. Thus there is concern protease inhibitors may adversely effect CTL epitope processing, but this paper indicates that processing is not inhibited at therapeutically relevant concentrations of RTV when the proteasome is functioning in an intracellular context.
- RTV did not alter the presentation two RT A2 epitopes processed by distinct pathways: ILKEPVHGV, generated by the constitutive proteasome containing the MB1 beta subunit, and VIYQYMDDL which is dependent on IFNgamma induction of LMP7 which replaces MB1 in the immunoproteasome, and is destroyed by MB1 in the constitutive proteasome.
- RTV did not inhibit the processing and assembly of HLA-B35 or -A2, which are assembled with a rapid and moderate time course, respectively, or of HLA-A3, -B27 and -B39.

HXB2 Location RT (179-187)

**Author Location** Pol (334–)

Epitope VIYQYMDDL

Epitope name Pol334

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

• HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.

- This epitope was one of the previously identified HLA-A2 epitopes studied. Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, proba-
- 1/17 HIV-infected HLA-A2+ people in this study recognized this epitope.

HXB2 Location RT (179–187)
Author Location Pol (334–342)
Epitope VIYQYMDDL
Immunogen HIV-1 infection
Species (MHC) human (A2)
Donor MHC A02, B35, Bw62

Assay type proliferation, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, memory cells, immune dysfunction

References Gamberg et al. 2004a

HAART restores HIV specific immunity after advanced infection by increase of CD4+ and CD8+ T cell numbers after supression of viral replication. However, HIV specific CTLs emerged only with detectable viral replication breakthroughs and were short-lived while CD4+ T-cell responses remained compromised, suggesting failure of generating stable CD8+ memory T-cells in the absence of HIV-specific T-helper responses.

HXB2 Location RT (179–187) Author Location RT (179–187) Epitope VIYQYMDDL Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2) Country Canada.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** HAART, ART, immunotherapy, variant crossrecognition or cross-neutralization

References Mason et al. 2004

- Accumulation of specific antiretroviral drug-resistance mutations in Pol gene was shown to sustain and even enhance the antigenicity and immunogenicity of HIV-1 CTL epitopes in this region. Several different patterns of cross-reactivity and selective recognition of wild-type and variant epitopes were found.
- VIcQYMDDL, VIYQYvDDL and VIcQYvDDL variants are detected due to appearence of Y181C and M184V resistance mutations. The double mutant was the only form recognized in one A02 treated individual, the epitope was not recognized in another.

HXB2 Location RT (179–187)
Author Location RT Pol (334–342)
Epitope VIYQYNDDL
Immunogen HIV-1 infection
Species (MHC) human (A2)
Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/19 patients recognized this epitope.

HXB2 Location RT (179–187) Author Location Pol Epitope VIYQMDDL

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2)

**Donor MHC** A\*02, A\*30, B\*4402, B\*15 **Assay type** Tetramer binding, T-cell Elispot

**Keywords** HIV exposed persistently seronegative (HEPS)

References Missale et al. 2004

- HIV-specific T-cell response was tested in patients exposed to blood from a patient with highly replicating HIV; these patients were nosocomially infected with HBV, but uninfected with HIV. HIV-specific T-cell responses were directed to structural and non-structural HIV proteins in 2 patients, suggesting that the virus replicated in these patients sufficiently to prime a cell-mediated immune response that protected them from HIV infection.
- This patient responded to 4/8 HIV epitopes tested in an IFNgamma EliSpot assay or tetramer assay. Responses were detected 8 and 28 weeks after exposure. No response to this epitope was detected by IFNgamma EliSpot, but a response was detected by tetramer staining.

HXB2 Location RT (179–187) Author Location RT (179–187) Epitope VIYQYMDDL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2) Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

 The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized in chronic infection.

HXB2 Location RT (179–187)
Author Location RT (179–187 HXB2)
Epitope VIYQYMDDL
Epitope name 51F
Subtype B
Immunogen vaccine

Vector/Type: DNA Strain: multiple epitope immunogen HIV component: p17/p24 Gag,

Pol Adjuvant: IL-12

**Species (MHC)** transgenic mouse (A2)

Assay type Cytokine production, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-specific epitope characteristics, vaccine antigen design

References Bolesta et al. 2005

- Immunization of transgenic mice with a codon-optimized hGagp17p24-Polp51 DNA plasmid, consisting of clusters of highly conserved CTL epitopes presented by multiple MHC class I alleles, induced 2- to 5-fold higher CD8+ T-cell responses than the corresponding full-length proteins. The modified proteins had the ribosomal frameshift deleted, as well as the potentially immunosuppressive p15, and protease and integrase. This correlated with higher protection against challenge with Gag and Pol expressing recombinant vaccinia virus. Mice immunized with the hGagp17p24-Polp51 also showed an elevated level of type 1 cytokine production as well as an increased titer of p24- and RT-specific IgG2 antibody responses.
- This was 1 of 4 A2 gag/pol epitopes tested. Transgenic mice immunized with the deleted construct induced more potent EliSpot reactions to this epitope than those immunized with full length Gag/Pol.

**HXB2 Location** RT (179–187)

Author Location RT (346–354)

Epitope VIYQYMDDL

Epitope name VL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Germany.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** HAART, ART, optimal epitope **References** Schmitt-Haendle *et al.* 2005

CTL responses to 3 HLA-A2-restricted epitopes were investigated in 51 HIV-1 infected HLA-A2+ individuals. The most prevalent response was seen for IV9, followed by SL9. The VL9 epitope was not recognized.

**HXB2 Location** RT (179–187)

**Author Location** Pol (subtype B)

Epitope VIYQYMMDL

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2, A\*0202)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among A, B and D clade viruses.

**HXB2 Location** RT (180–189)

**Author Location RT (LAI)** 

Epitope IYQYMDDLYV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**References** Menendez-Arias *et al.* 1998; van der Burg *et al.* 1997

- Recognized by CTL from a progressor, spans important RT functional domain.
- A previous study determined that this was an epitope recognized by a long-term survivor.

**HXB2 Location** RT (181–189)

Author Location RT (181–189 LAI)

Epitope YQYMDDLYV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** binding affinity, computational epitope prediction

References Samri et al. 2000

- This epitope contains the mutation M184V, a frequent mutation induced by nucleoside reverse transcriptase inhibitors.
- High levels of recognition by ELISPOT were observed for zidovudine induced mutation YQYVDDLYV and for the wildtype peptide YQYMDDLYV in patient 250#0 (HLA-A\*0201), but neither were recognized by patient 201#5 (also HLA-A\*0201)
- Both the wild-type and the mutated peptide were computer predicted to have a high binding affinity for A2 (http://bimas.dcrt.nih.gov/molbio/hla\_bind)

**HXB2 Location** RT (192–201)

Author Location RT (192-201)

Epitope DLEIGQHRTK

Immunogen HIV-1 infection

Species (MHC) human (A3)

References Haas et al. 1998

- Of 98 patients in cross-sectional analysis, 78% had CTL against pol RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)
- New clusters of epitopes were defined utilizing different HLA molecules.

**HXB2 Location** RT (192–201)

**Author Location** Pol (192–201)

Epitope DLEIGQHRTK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 7 epitopes from this individual that varied over time, although the internal mutations did not become fixed. The dleMgqhrtk variant arose at late time points.

HXB2 Location RT (192–216) Author Location RT (359–383 HXB2)

Epitope DLEIGQHRTKIEELRQHLLRWGLTT

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B60)

**References** Menendez-Arias *et al.* 1998; Walker *et al.* 1989

 One of five epitopes defined for RT-specific CTL clones in this study.

HXB2 Location RT (192–216) Author Location RT (191–215)

Epitope DLEIGQHRTKIEELRQHLLRWGFTT

Immunogen HIV-1 infection
Species (MHC) human (polyclonal)
Keywords HAART, ART, escape

References Haas et al. 1997; Menendez-Arias et al. 1998

 Polyclonal CTL recognition switched from RT 191-215 to RT 514-524 when AZT therapy selected for the resistance mutation, and presumably the escape variant, RT T215Y.

**HXB2 Location** RT (198–212) **Author Location** RT (SF2)

Epitope HRTKIEELRQHLLRW Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.
- The response to the peptide was CD8 dependent, but the HLA presenting molecule and optimal epitope were not determined.

**HXB2 Location** RT (201–209)

**Author Location** RT (201–209)

**Epitope** KIEELRQHL **Immunogen** HIV-1 infection

Species (MHC) human (A2)

References Haas et al. 1998

- Of 98 patients in cross-sectional analysis, 78% had CTL against pol RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)
- New clusters of epitopes were defined utilizing different HLA molecules.

**HXB2 Location** RT (201–210)

Author Location Pol

Epitope KIEELRQHLL

Immunogen

Species (MHC) human (B58)

References De Groot et al. 2001

- The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.
- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes could stimulate IFNγ production in an ELISPOT assay.

- KIEELRQHLL was newly identified as a HLA-B58 epitope in this study, it had been previously shown to be presented by HLA-A2 and Bw60.
- KIEELRQHLL did not bind detectably to B7.

**HXB2 Location** RT (202-210)

Author Location RT (202–210 LAI)

Epitope IEELRQHLL

Subtype B

Immunogen

Species (MHC) human (B\*4001)

Keywords optimal epitope

References Altfeld et al. 2000; Frahm et al. 2007

• C. Brander notes this is a B\*4001 epitope.

HXB2 Location RT (202–210)

Author Location RT (SF2)

Epitope IEELRQHLL

Immunogen HIV-1 infection Species (MHC) human (B60)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B60+ individuals that had a CTL response to this epitope broken down by group: 0/2 group 1, 1/1 group 2, and 0/0 group 3.

**HXB2 Location** RT (202–210)

**Author Location RT (SF2)** 

Epitope IEELRQHLL

Immunogen HIV-1 infection

Species (MHC) human (B60)

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes.
- B60 is present in 10-20% of the Caucasoid and very common in Asian populations.

**HXB2 Location** RT (202-210)

**Author Location RT** 

Epitope IEELRQHLL

Epitope name IL9

Immunogen HIV-1 infection

Species (MHC) human (B60)

**Donor MHC** A2, A24, B38, B60, Cw2, Cw12

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** HAART, ART, supervised treatment interruptions (STI), early treatment

References Montefiori et al. 2003

• HIV-1 + patient AC10 underwent early HAART treatment, which was discontinued 1.5 years later. At this timepoint potent NAb responses against autologous virus were detected. Treatment interruption initially induced weak CD8+ responses directed against 5 epitopes. By days 873d and 923d the CTL response had broadened to target 22 epitopes; of these six were fully characterized. Eventually the virus escaped the NAb response, but escape was not accompanied by a rise in viral load, and the authors suggest the virus was contained by the CTL response.

HXB2 Location RT (202–210)
Author Location RT (202–210)
Epitope IEELRQHLL
Immunogen HIV-1 infection
Species (MHC) human (B60, B61)
Keywords immunodominance
References Day et al. 2001

- No immunodominant responses were detected to five B61restricted epitopes tested.
- All five B60-restricted epitopes were reactive in another subject, and the B60-restricted responses together contributed over onethird of the total CTL response.

**HXB2 Location** RT (203–211)

**Author Location RT** 

Epitope EELRQHLLR Immunogen HIV-1 infection Species (MHC) human (B44)

Donor MHC A\*3101, A68, B\*4403, B51

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Arnedo-Valero et al. 2004

- T cell immune responses following STI were monitored in two chronically HIV-1 infected partners (A and B) who had contracted HIV-1 during 1992. Both were heterozygous for the CCR5 delta32, and had different HLAs and treatment histories. STI induced strong transitory CD4+ and CD8+ T cell responses in both patients. Patient A displayed broad CD8+ T cell responses directed against Env, Pol, Gag, and Nef HIV-1 antigens. CTL responses in patient B were mainly directed against two epitopes: Gag(p24)NANPDSKTI and Pol(RT)EELRQHLLRW.
- Despite the host differences, both patients had similar dynamics of vial evolution and CD4+ T-cells, suggesting that good immune reponses to STI may be more related to the virus than host characteristics in these cases.

**HXB2 Location** RT (203–212)

**Author Location** RT (LAI)

Epitope EELRQHLLRW

Subtype B

**Immunogen** HIV-1 infection **Species** (MHC) human (B44)

**References** Menendez-Arias *et al.* 1998; van der Burg *et al.* 1997

• The only epitope recognized by CTL from a long-term survivor in two samples taken six years apart.

 Recognized by CTL from a progressor, EILKEPVGHGV and TWETWWTEYW were also recognized.

**HXB2 Location** RT (203–212)

**Author Location** RT (203–212)

Epitope EELRQHLLRW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B44)

Country Canada.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

**Keywords** HAART, ART, immunotherapy, variant crossrecognition or cross-neutralization

References Mason et al. 2004

- Accumulation of specific antiretroviral drug-resistance mutations in Pol gene was shown to sustain and even enhance the antigenicity and immunogenicity of HIV-1 CTL epitopes in this region. Several different patterns of cross-reactivity and selective recognition of wild-type and variant epitopes were found.
- EELRQHLwRW variant is detected due to appearence of L210W resistance mutation. The in this case, the wild-type epitope was preferentially recognized relative to the L210W variant.

**HXB2 Location** RT (203–212)

**Author Location** RT Pol (358–367)

Epitope EELRQHLLRW

Immunogen HIV-1 infection

Species (MHC) human (B44)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 9/11 patients recognized this epitope; of three B\*44 eptiopes tested, this was the only one that was recognized by more than 2/11 patients.

**HXB2 Location** RT (203–212)

**Author Location RT** 

Epitope EELRQHLLRW

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B44)

**Donor MHC** A01, A03, B39, B44, Cw4, Cw6

Assay type T-cell Elispot

Keywords HIV exposed persistently seronegative

(HEPS)

References Missale et al. 2004

- HIV-specific T-cell response was tested in patients exposed to blood from a patient with highly replicating HIV; these patients were nosocomially infected with HBV, but uninfected with HIV. HIV-specific T-cell responses were directed to structural and non-structural HIV proteins in 2 patients, suggesting that the virus replicated in these patients sufficiently to prime a cell-mediated immune response that protected them from HIV infection.
- This patient responded to 3/11 HIV epitopes tested in an IFNgamma EliSpot assay. Responses were detected 16 and 20 weeks after exposure, but were lost by week 80.

**HXB2 Location** RT (203-212)

**Author Location** p24

Epitope EELRQHLLRW Immunogen HIV-1 infection Species (MHC) human (B44)

**Donor MHC** A01, A32, B\*1410, B15; A\*3101, A68,

B\*4403, B51

Country Spain.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI)

References Arnedo-Valero et al. 2004

- T cell immune responses following STI were monitored in two chronically HIV-1 infected partners (A and B) who had contracted HIV-1 during 1992. STI induced strong transitory CD4+ and CD8+ T cell responses in both patients. The viruses remained very closely related over 10 years, despite the two individuals having different HLA types; the authors suggest the maintained similarity does not support a strong role for HLA driven HIV diversity as has been claimed in Moore et al. (Science 2002).
- During the second treatment stop, patient A developed a strong proliferative response to p24, and multiple strong CD8+ T cell repsonses to Env, Pol, Gag and Nef. This patient was able to control viral load for two years follow up without therapy. Patient B developed a very weak CD4+ T cell response against p24 during breaks in therapy, and had CD8+ responses to two epitopes. Patient A: A01, A32, B\*1410, B15; Patient B: A\*3101, A68, B\*4403, B51.

**HXB2 Location** RT (203–212)

Author Location RT (203–212)

Epitope EELRQHLLRW

**Immunogen** HIV-1 infection **Species** (MHC) human (B44)

Constant Constant

Country Canada.

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords mimics

References Mason et al. 2005

 CTL responses against the IP-30 signal peptide associated with autoimmunity were shown to be elicited by stimulation of PBMCs from HIV-1 infected individuals with HIV protease peptide 76-84. In vitro stimulation with HIV PR 76-84 or the IP-30 signal peptide was shown to activate a comparable population of cross-reactive effector cells. None of the peptides activated CTL in non-HIV-infected individuals. IP-30 signal peptide was shown to have lower avidity T-cell interactions than the HIV peptide.

 As a control, responses to A2-restricted HIV epitopes ALVE-ICTEM, EELRQHLLRW, and LSPRTLNAW were shown not to give IP-30 responses.

**HXB2 Location** RT (204–212)

**Author Location** RT (204–212 HXB2)

Epitope ELRQHLLRW

Epitope name EW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2501)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- This is a newly defined epitope. Position 8 in the epitope had potentially experienced positive selection. ELRQHLLkW escape variant was found.

**HXB2 Location** RT (209–220)

**Author Location** RT (209–220 MN)

**Epitope** LLRWGLTTPDKK

Subtype B

Immunogen vaccine

Vector/Type: DNA, polyepitope Strain: B clade MN HIV component: gp120, Protease, RT Adjuvant: Incomplete Freund's Adju-

vant (IFA)

**Species (MHC)** humanized mouse (A\*0201)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance, im-

munotherapy

References Isaguliants et al. 2004

- Immunization of HLA-A\*0201-transgenic mice with synthetic genes encoding clusters of human A\*0201 CTL epitopes located at the sites of drug resistance mutations, induced RTspecific cellular responses indicating the immunogenicity of these constructs. This vaccine strategy may be a first step towards a therapeutic vaccine against drug-resistant strains.
- This was one of five HLA-A\*0201 epitopes from the RT or protease proteins that was included in the polyepitope vaccine. When the transgenic HLA A\*0202 mice were vaccinated with the polyepitope construct or with a mixture of RT peptides, a sustained low level CD8+ T-cell gamma IFN response was observed, in contrast to when an intact RT gene was used for vaccination.

Author Location RT (209–220)

Author Location RT (209–220)

Epitope LLRWGLTTPDKK

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Haas et al. 1998

- Of 98 patients in cross-sectional analysis, 78% had CTL against pol – RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)
- New clusters of epitopes were defined utilizing different HLA molecules.

Author Location RT (210–220)
Author Location Pol (209–220)
Epitope LRWGFCTPDKK
Immunogen HIV-1 infection
Species (MHC) human (B57)

Donor MHC A2, B44, B57; A2, A29, B57, B62

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** HAART, ART, cross-presentation by different HLA, characterizing CD8+ T cells

References Mason & Grant 2005

- The common form of this epitope, LRWGFTTPDKK is weakly recognized in the context of HLA-A2, and it encompasses several common antiretroviral escape mutations. Responses were tested in 2 siblings.
- T215Y then Y215C antiretroviral therapy-associated mutations within the epitope induced a strong reaction, but changed the restriction of the epitope to HLA-B57. This mutation is thus suggested to potentially enhance CD8 T-cell recognition of HIV.

**HXB2 Location** RT (214–223)

**Author Location** Pol

Epitope FTTPDKKHQK

Epitope name 1267
Subtype multiple
Immunogen HIV-1 infection
Species (MHC) human (A11, A68)

**Donor MHC** A11, A68, B42, B45, Cw16, Cw17

**Country** United States. **Assay type** T-cell Elispot

**Keywords** binding affinity, supertype, computational epitope prediction, cross-presentation by differ-

ent HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for FTTPDKKHQK:36% Supertype epitope binding to A68 and A11.

**HXB2 Location** RT (215-224)

**Author Location** Pol

Epitope TTPDKKHQKE

Subtype multiple
Immunogen HIV-1 infection
Species (MHC) human (A11)

 $\textbf{Donor MHC} \ \, \text{A11, A68, B42, B45, Cw16, Cw17; A01,} \\$ 

A68, B15, B40, Cw03

Country United States.
Assay type T-cell Elispot

**Keywords** binding affinity, supertype, computational epi-

tope prediction, cross-presentation by differ-

ent HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for TTPDKKHQKE:60% Supertype epitope binding to A68.

**HXB2 Location** RT (218–235)

**Author Location** (C consensus)

Epitope DKKHQKEPPFLWMGYELH

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*1510)

Country South Africa.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (240–257)

**Author Location** (C consensus)

Epitope TVQPIQLPEKDSWTVNDI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (240–257)

**Author Location** RT (240–257 HXB2)

Epitope TVQPIVLPEKDSWTVNDI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%).
   p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- Responses to this peptide were detected in 16% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** RT (243-252)

**Author Location** RT (LAI)

Epitope PIVLPEKDSW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

**References** Menendez-Arias *et al.* 1998; van der Burg *et al.* 1997

 Recognized by CTL from a progressor and a long-term survivor, KITTESIVIW was also recognized.

**HXB2 Location** RT (243–252)

**Author Location** RT (LAI)

Epitope PIVLPEKDSW

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*5701)

**Keywords** binding affinity, escape

**References** Menendez-Arias *et al.* 1998; van der Burg *et al.* 1997

Recognized by CTL from long-term survivor, whose CTL response persisted for more than 10 years – the substitution V3M reduced affinity but was well recognized, on the other hand V3T and D8G did not reduce affinity, but abrogated CTL response.

**HXB2 Location** RT (243–252)

**Author Location** RT (410–419)

Epitope PIVLPEKDSW

Epitope name PIV

Immunogen HIV-1 infection Species (MHC) human (B57)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- None of the 8 study subjects recognized this epitope but none were HLA B57+

**HXB2 Location** RT (243-252)

**Author Location RT** 

Epitope PIVLPEKDSW

Epitope name PIV

Immunogen HIV-1 infection Species (MHC) human (B57)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius *et al.* [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there
  was no correlation between CTL responses with viral rebound
  rates, plateau viral loads, or clearance rates.

**HXB2 Location** RT (243–252)

**Author Location** RT Pol (398–407)

Epitope PIVLPEKDSW

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 6/7 patients recognized this epitope.

**HXB2 Location** RT (244–252)

Author Location RT (399-407)

Epitope IVLPEKDSW

Immunogen

Species (MHC) human (B\*5701)

Keywords optimal epitope

References Frahm et al. 2007

- Subtype of B57 not determined.
- C. Brander notes this is a B\*5701 epitope.

**HXB2 Location** RT (244–252)

**Author Location** RT (244–252 LAI)

**Epitope IVLPEKDSW** 

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5701, B\*5801)

Keywords binding affinity, rate of progression, escape

References Klein et al. 1998

- This peptide was defined as the optimal epitope.
- B57 has been associated with long-term non-progression in the Amsterdam cohort.
- The most pronounced CTL responses in HLA B\*5701 LTS were to RT and Gag.
- B57 restricted CTL responses are targeted at multiple proteins, but one LTS had a response that was dominated by reactivity to the epitope two variants were found in this LTS: ITLPEKESW, which bound to B\*5701 with similar affinity as the index peptide but was an escape mutant that was not recognized by CTL, and IMLPEKDSW, which bound to B\*5701 with reduced affinity but could still be recognized.
- In an additional HIV+ LTS, only the variant IELPEKDSW was found, and this epitope was recognized by CTL but had less affinity for B\*5701 than the index peptide.
- This epitope was recognized in the context of both HLA-B\*5701 and B\*5801.

**HXB2 Location** RT (244–252)

**Author Location** Pol (244–252)

Epitope IVLPEKDSW

Immunogen HIV-1 infection

Species (MHC) human (B\*5801)

References Appay et al. 2000

- Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virus-specific CD8+ T cells produced IFN- $\gamma$  and MIP-1 $\beta$  with a distinct subset that failed to produce TNF- $\alpha$

**HXB2 Location** RT (244–252)

**Author Location** RT (399–407)

**Epitope IVLPEKDSW** 

Immunogen

Species (MHC) human (B57)

References van der Burg et al. 1997

**HXB2 Location** RT (244–252)

**Author Location** RT (244–252)

**Epitope** IVLPEKDSW **Immunogen** HIV-1 infection

Species (MHC) human (B57)

Keywords early-expressed proteins, kinetics

References Guillon et al. 2002b

 An early-expressed Nef protein was modified to contain Env and Pol epitopes to enable the study the effect of expression kinetics on CTL mediated suppression of replication. The "EpiNef" construct was inserted into a recombinant vaccinia virus which was used to infect a target cell line; the target cells were lysed by CTL clones specific for the Env and Pol epitopes indicating that they were properly processed.

**HXB2 Location** RT (244–252)

**Author Location** RT (244–252 ACH320.2A.2.1)

**Epitope IVLPEKDSW** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) (B57)

**Keywords** acute/early infection, early-expressed proteins, kinetics

References van Baalen et al. 2002

• Tat, Rev and Nef are the first HIV proteins expressed upon acute infection of T-cells (< 6 hours), and RT is not expressed until after 24 hours. The B14-restricted Rev-SAEPVPLQL specific CD8 T-cell clone TCC108, and the B57-restricted RT-IVLPEKDSW specific CD8 T-cell clone TCL1C11 were co-incubated with CD4+ cultures innoculated with HIV-1 at low MOI. Co-incubation with the Rev-specific CTL resulted in two logs less HIV-1 production in ten days of culture. When the RT epitope was cloned into the Nef gene of the infecting strain, another early expressed protein, it proved as effective as the Rev epitope at inhibiting viral production. A mathematical model of CTL-target interactions suggest early proteins are important for vaccine design.

**HXB2 Location** RT (244–252)

**Author Location** Pol

Epitope IVLPEKDSW

Epitope name IW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 2, was found in the most polymorphic residue in the epitope. This was shared between clades B and C. The position 2 mutation was significantly more common among persons expressing HLA-B57.

**HXB2 Location** RT (244–252)

**Author Location** 

**Epitope** IVLPEKDSW

Immunogen HIV-1 infection

Species (MHC) human (B57, B\*5801)

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

Keywords responses in children, mother-to-infant trans-

mission, escape

References Feeney et al. 2005

• Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.

**HXB2 Location** RT (245–252)

Author Location Pol

**Epitope IVPEKDSW** Immunogen HIV-1 infection Species (MHC) human (B57)

References Kostense et al. 2001

- HLA tetramers to six epitopes were used to study HLA-A2, B8 and B57 CTL in 54 patients - HIV-specific tetramer positive cells were inversely correlated with viral load in patients with high CD4, but in patients with CD4 T-cells below 400 high tetramer frequencies were found despite high viral load.
- · Most patients have high levels of HIV-specific T-cell expansions, but many of these cells aren't functional.
- In 15 of the patients, the proportion of IFN gamma producing tetramer cells correlated with AIDS-free survival.

**HXB2 Location** RT (248–264)

Author Location (C consensus)

Epitope EKDSWTVNDIQKLVGKL

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*0205)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (259-267)

**Author Location** Pol

Epitope KLVGKLNWA Immunogen HIV-1 infection **Species (MHC)** human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.

- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind four of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).
- Tetramer staining with A2, beta2microglobulin, and either SLYNTVATL, KLVGKLNWA, or LTFGWCFKL revealed that tetramers detected more HIV-specific sells in LTNP than in progressors, activated effector cells were the minority population, and ELISPOT correlated better with the effector cell subpopulation than the total tetramer stained population.

**HXB2 Location** RT (260–271)

**Author Location** RT (415–426 IIIB)

Epitope LVGKLNWASQIY

Immunogen HIV-1 infection

Species (MHC) human (B\*1501)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is a B\*1501 epitope.

**HXB2 Location** RT (260–271)

**Author Location** Pol (260–271)

Epitope LVGKLNWASQIY

Immunogen HIV-1 infection

Species (MHC) human (B15)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

**Country** Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 7 epitopes from this individual that varied over time, although the internal mutations did not become fixed. lvgkXnwasqiy variants arose at late time points

HXB2 Location RT (260-271)

**Author Location** RT (415–426 IIIB)

Epitope LVGKLNWASQIY

Immunogen HIV-1 infection

Species (MHC) human (B62)

References Brander & Walker 1996; Menendez-Arias

et al. 1998

• P. Johnson, pers. comm.

**HXB2 Location** RT (260–271)

**Author Location** RT (260–271)

Epitope LVGKLNWASQIY

Immunogen HIV-1 infection

Species (MHC) human (B62)

Keywords immunodominance References Day et al. 2001

 No immunodominant responses were detected to four B62 In subject 199 four additional A\*3002 epitopes were identified. restricted epitopes tested.

**HXB2 Location** RT (260-271) Author Location Pol (415-426) Epitope LVGKLNWASQIY Epitope name LY12 Subtype B

Immunogen HIV-1 infection Species (MHC) human (B62)

**Donor MHC** A\*01, A\*11, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords escape, characterizing CD8+ T cells, optimal epitope

References Koibuchi et al. 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- The autologous form of the epitope, LVGKLNWASQIY, matched the B consensus throughout the 5-year period of study, with 1 rare variant at the first time point: LVGKLNWASQIh.

**HXB2 Location** RT (263–271) **Author Location** RT (263–271 LAI) Epitope KLNWASOIY Subtype B

**Immunogen** 

Species (MHC) human (A\*3002)

Keywords optimal epitope

References Frahm et al. 2007; Goulder et al. 2001a

• C. Brander notes this is an A\*3002 epitope.

**HXB2 Location** RT (263-271)

**Author Location RT** 

Epitope KLNWASQIY Epitope name KY9 (RT-35) Immunogen HIV-1 infection Species (MHC) human (A\*3002)

References Goulder et al. 2001a

- HLA-A\*3002 is very common in African populations, 50% of Zimbabweans express HLA-A30, 44% in African Zulu, so five new HIV epitopes were characterized that are presented by this HLA molecule.
- · A rapid method was developed combining ELISPOT with intracellular IFN- $\gamma$  staining of PBMCs to map optimal epitopes, then HLA presenting molecules were defined - this method was completed within 48 to 72 hours of receipt of blood.
- Subject 199 (HLA • Two individuals were studied: A\*0201/\*3002 B\*4402/51 Cw2/5), a Caucasian, and Subject 6007 (HLA A\*3002/ B53/\*5801 Cw4/7) an African-Caribbean.
- In both HLA-A\*3002 individuals the response to RSLYNT-VATLY was dominant.

- Three quantitative assays, ELISPOT, precursor frequency and chromium release, confirmed a hierarchy of response: RY11 (p17) > KY9 (gp41) > KY9 (RT-53) > IY9 (gp41)

**HXB2 Location** RT (263-271)

Author Location (C consensus)

Epitope KLNWASQIY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** RT (263–271)

Author Location (C consensus)

**Epitope** KLNWASQIY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- KLNWASQIY is an optimal epitope.

**HXB2 Location** RT (263-271)

**Author Location RT** 

Epitope KLNWASQIY

**Epitope name** A30-KY11(RT)

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A30)

**Donor MHC** A30, A32, B18, B27

Keywords HAART, ART, supervised treatment interrup-

tions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals. Patient D displayed the greatest response to B27-KK10 (p24), and also responded to A30-RY11(p17), A32-PW10(RT), A30-KY11(RT), A32-RW10(gp120), and B18-YY9(Nef).

 $\textbf{HXB2 Location} \ \ \mathsf{RT} \ (266\text{--}285)$ 

Author Location Pol (421–440)

Epitope WASQIYPGIKVRQLCKLLRG

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons

References Novitsky *et al.* 2002
• HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from

between 55 and 64 subjects for each HIV protein.

- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** RT (268–282)

**Author Location RT (SF2)** 

Epitope SQIYPGIKVRQLCKL

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2001a

- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- RT peptides SQIYPGIKVRQLCKL and WKG-SPAIFQSSMTKI were recognized.

**HXB2 Location** RT (269–277) **Author Location** (LAI)

Epitope QIYPGIKVR

Subtype B

Immunogen

Species (MHC) human (A\*0301)

Keywords optimal epitope

References Altfeld 2000; Frahm et al. 2007

**HXB2** Location RT (269–277)

Author Location RT (269–277)

Epitope QIYPGIKVR

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Assay type Other

Keywords HLA associated polymorphism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- QIYPGIKVR was a previously defined A\*0301 presented epitope that encompassed an A\*03- associated polymorphism, QIYPGIKVrlQ, in the last position. This epitope was embedded in a previously determined CTL immunoreactive region.

**HXB2 Location** RT (269–277)

**Author Location** Pol (424–432)

Epitope QIYAGIKVK

Subtype B, CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A\*1101)

Keywords binding affinity, subtype comparisons

References Fukada et al. 2002

- binding affinity, inter-clade comparisons.
- Counterparts for eight known clade B HLA A\*1101 epitopes were generated for clade E (CRF01). Three epitopes, identical among clade A-E, were cross-reactive and recognized by clade E infected individuals. The clade E and B analogs to three more HLA A\*1101 epitopes was recognized in a clade specific manner. Two other HLA A\*1101 clade B defined epitopes were found not to have stimulated a response in clade E infected individuals.
- QIYAGIKVK is commonly found in viruses representing subtypes A, B and E. It was strongly recognized by CTL from 1/5 B clade infected Japanese subjects, and 5/7 E clade infected Thai subjects.
- QIYAGIKVK had the highest A\*1101 binding affinity, but qiyagikvR and qiyPgikvR (the most common C and D clade variant both bound to A\*1101). QIYAGIKVK and qiyagikvR were both cross-presented by a clone from a B clade infection, but qiyPgikvR was not.

**HXB2 Location** RT (269–277)

Author Location (B consensus)

Epitope QIYAGIKVK

Epitope name QVK9

Immunogen HIV-1 infection

Species (MHC) human (A11)

**Donor MHC** A02, A11, B18, B44, Cw5, Cw12

Country United States.

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

 $\textbf{Keywords} \ \ \text{assay standardization/improvement, memory}$ 

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** RT (269-277)

**Author Location** Pol (425–433)

Epitope QIYAGIKVK

Epitope name QK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Flow cytometric T-cell cytokine

assay

Keywords optimal epitope

References Allen et al. 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For one of the escape variants, a novel CD8 T-cell response equal in magnitude to the wild type, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wild type.
- This epitope did not vary.

**HXB2 Location** RT (269–277)

Author Location Pol (425-433)

Epitope QIYAGIKVK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.

 This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** RT (269–277)

**Author Location** RT (269–277)

Epitope QIYPGIKVR

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- All patients recognized at least 1 A3 epitope, up to 8 A3 epitopes, but none was clearly dominant.

**HXB2 Location** RT (269–277)

**Author Location** RT (424–432)

Epitope QIYPGIKVR

Epitope name A3-OR9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A3, B7, Cw7

Keywords dynamics, supervised treatment interruptions

(STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 1/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 4/7 individuals began to have detectable responses to this epitope after STI.

**HXB2 Location** RT (269–277)

Author Location RT (269-277)

Epitope QIYAGIKVK

Epitope name A3-QR9 Pol

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Assay type CD8 T-cell Elispot - IFNγ

Keywords supervised treatment interruptions (STI), es-

cape, early treatment, superinfection

References Altfeld et al. 2002a

- An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.
- The second infecting strain had the variant qiyagikvR. The initial CTL response to both variants was strong but eventually declined, particularly to the variant in the second strain. .

**HXB2 Location** RT (269–277) **Author Location** RT (269–277) Epitope QIYPGIKVR Immunogen HIV-1 infection Species (MHC) human (A3)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4, DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- · Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

**HXB2 Location** RT (269–277)

**Author Location** Pol

Epitope QIYPGIKVK

Epitope name QK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- · Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 9, qiypgikvR, was found not to correspond to the most polymorphic residues in the epitope.

**HXB2 Location** RT (271-279)

**Author Location (LAI)** 

Epitope YPGIKVRQL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*4201 epitope.

**HXB2 Location** RT (271–279)

Author Location (C consensus)

Epitope YPGIKVRQL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** RT (271–279)

**Author Location** (C consensus)

Epitope YPIGKVRQL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the L9 residue of YPIGKVRQL are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** RT (271–279)

Author Location RT (271-279)

Epitope YPGIKVRQL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Assay type Other

Keywords HLA associated polymorphism

References Boutwell & Essex 2007

- HIV-1 subtype C proteins were analysed and 94 HLA- Detection of CTL escape mutants in the mother was associated associated amino acid polymorphisms were found. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 88% of CTL immunodominant regions had same HLA-associations as found polymorphisms. In 19 studied CTL epitopes, the epitope-restricting allele was same as the polymorphism-restricting allele for 7, and in 12 the alleles differed by specificity.
- YPGIKVRQL was a previously defined B\*4201 presented epitope that encompassed an B\*42 associated polymorphism, YpGIKVRQL,in the second position. This epitope was found embedded in a previously determined immunoreactive region.

**HXB2 Location** RT (271–279)

**Author Location** (C consensus)

Epitope YPIGKVRQL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4202)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the L9 residue of YPIGKVRQL are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** RT (271–279)

**Author Location** RT (438–446 IIIB)

Epitope YPGIKVRQL Immunogen HIV-1 infection

Species (MHC) human (B42) Keywords responses in children, mother-to-infant transmission

References Menendez-Arias et al. 1998; Wilson et al.

- YAGIKVRQL and YPGIKVKQL are naturally occurring variants that are both reactive.
- YHKIKVRQL is a naturally occurring variant that has not been
- · Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.

**HXB2 Location** RT (271–279)

Author Location Pol (438–446 IIIB)

Epitope YPGIKVRQL Immunogen HIV-1 infection

Species (MHC) human (B42) Keywords mother-to-infant transmission, escape

References Wilson et al. 1999a

· This study describes maternal CTL responses in the context of mother-to-infant transmission.

- with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- An additional variant that gave a positive CTL response: YPGIKVKQL, YAGIKVRQL.
- YHGIKVRQL was an escape mutant.

**HXB2 Location** RT (293-301)

**Author Location** RT (448–456 SF2)

**Epitope IPLTEEAEL** Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

References Menendez-Arias et al. 1998; Tomiyama et al.

- A CTL clone responsive to this epitope was obtained.
- Only 1/7 B35-positive individuals had a CTL response to this epitope.
- An E to K substitution at position 5 abrogates specific lysis, but not binding to B\*3501.
- An I to V substitution at position 1, P to Q at position 2, and E to K at 5, abrogates specific lysis and binding to B\*3501.
- An I to V substitution at position 1 did not alter reactivity.
- Reviewed in Menendez-Arias et al. [1998], this epitope lies in the thumb region of RT.

**HXB2 Location** RT (293-301)

Author Location Pol (HXB2, LAI)

**Epitope IPLTEEAEL** 

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B\*3501)

**Donor MHC** A\*2402, A\*2601, B\*3501, B\*5101

Country Japan.

Assay type Cytokine production, Tetramer binding, Chromium-release assay

Keywords binding affinity, kinetics, TCR usage, characterizing CD8+ T cells, immune dysfunction

References Ueno et al. 2004b

- Two different clonotypes of CD8+ T-cells with specificity for this epitope were isolated from a chronic HIV+ patient. The clonotype with the relatively high affinity TCR had no cytolytic activity, cytokine production or proliferation in response to HIV-infected cells, while the moderate affinity clonotype had strong reactions. More than 3-fold increased duration in tetramer 1/2 life was observed with the defective clonotype. The TCRs from the two clonotypes preserved the phenotype when transduced into primary CD8+ T cells, suggesting the TCR with higher affinity was directly associated with impaired T-cell reactivity of the cells.
- The high affinity impared TCR was Valpha1.1/Vbeta13.3, the moderate affinity active TCR was Valpha12.1/Vbeta5.6.

**HXB2 Location** RT (293–301)

Author Location Pol (HXB2, LAI)

**Epitope IPLTEEAEL** 

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*3501, B\*1501)

**Donor MHC** A\*2402, A\*2601, B\*3501, B\*5101

Country Japan.

Assay type Cytokine production, Tetramer binding, Chromium-release assay

**Keywords** binding affinity, cross-presentation by different HLA, immunotherapy, TCR usage, characterizing CD8+ T cells

References Ueno et al. 2004a

 This paper described the transduction of HIV specific clone TCR genes Valpha12.1/Vbeta5.6 into primary CD8+ T cells. Epitope fine specificity and appropriate effector functions were observed in the transduced cells, although functional avidity could change due to different densities of TCR on the surface of the transduced cells. No allogenic responses were detected. This methodology could have immunotherapeutic applications.

**HXB2 Location** RT (293-301)

Author Location Pol (448-456 SF2-24)

Epitope IPLTEEAEL
Epitope name HIV-B35-SF2-24
Immunogen HIV-1 infection

**Species (MHC)** human (B\*3501, B\*5101)

References Tomiyama et al. 2000b

- This epitope is naturally processed and presented by both HLA-B\*3501 and HLA-B\*5101 and is cross-recognized by a single CTL clone.
- IPLTEEAEL binds approximately four times more tightly to HLA-B\*3501 than HLA-B\*5101.

**HXB2 Location** RT (293-301)

**Author Location** Pol (489–456)

Epitope IPLTEEAEL

Immunogen HIV-1 infection

**Species (MHC)** human (B\*3501, B\*5301, B\*5101, B\*0702)

**Donor MHC** A24, A26, B35, B51, Cw3

**Keywords** supertype, cross-presentation by different HLA, TCR usage

References Ueno et al. 2002

The IPLTEEAEL epitope was known to be presented by both HLA-B\*3501 and -B\*5101 to a dual specific CTL clone. A single TCR complex bearing Valpha12.1 and Vβ5.6 was shown recognize the epitope in either HLA-B\*3501 and -B\*5101. Furthermore, this TCR also recognized the peptide presented by B\*5301 and B\*0702 in cytolytic CTL assays, demonstrating that this single TCR complex recognizes the same peptide presented by a range of HLA class I molecules.

**HXB2 Location** RT (293–301)

**Author Location (SF2)** 

Epitope IPLTEEAEL
Immunogen HIV-1 infection
Species (MHC) human (B35)
Keywords rate of progression
References Kawana *et al.* 1999

- HLA B35 is associated with rapid disease progression.
- The sequences of 9 previously described HIV-1 B35 CTL epitopes were obtained in 10 HLA B35+ and 19 HLA B35- individuals.
- 3/9 CTL epitopes had substitutions that were more common in B35+ individuals than in B35- individuals but this was one of the six that had no B35 associated pattern of mutation.

**HXB2 Location** RT (293–301)

Author Location RT (448–456 SF2)

**Epitope IPLTEEAEL** 

Immunogen HIV-1 infection

Species (MHC) human (B35, B51)

References Menendez-Arias et al. 1998; Shiga et al. 1996

- Binds HLA-B\*3501 and B\*5101.
- Reviewed in Menendez-Arias et al. [1998], this epitope lies in the thumb region of RT.

**HXB2 Location** RT (293–301)

Author Location Pol (447–455)

**Epitope IPLTEEAEL** 

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B51)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

• ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi

sex workers.

**HXB2 Location** RT (293–301)

Author Location RT (293-301)

**Epitope IPLTEEAEL** 

Immunogen HIV-1 infection

Species (MHC) human (B51)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DO7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

**HXB2 Location** RT (293-301)

Author Location RT Pol (286–294)

Epitope IPLTEEAEL

Epitope name IPL

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401, Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

DO6

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Tetramer binding,

CD4 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, immunodominance, escape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant eptiopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- The IPL epitope was found to be under positive selection for escape mutations and it was replaced by first variant between days 297 and 369, ipltGeael. This new variant was subsequently replaced by 2 further variants, that were even more resistent to CD8+ T cell recognition between days 369 and 635, ipltAeael and ipltVeael.

**HXB2 Location** RT (294–318)

**Author Location** RT (461–485 HXB2)

Epitope PLTEEAELELAENREILKEPVHGVY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**References** Menendez-Arias *et al.* 1998; Walker *et al.* 

• One of five epitopes defined for RT-specific CTL clones in this study.

**HXB2 Location** RT (298–312)

**Author Location** RT (291–305)

**Epitope** EAELELAENREILKE

**Epitope name** EAE

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401, Cw\*0702, DR17, DR15, DR51, DR52, DO2,

DQ6

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot

- IFNγ

**Keywords** rate of progression, escape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant epitopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This was one of 8 reactive peptides found not to vary over time. It was one of four epitopes that were not precisely defined.

**HXB2 Location** RT (302–319)

**Author Location** (C consensus)

Epitope ELAENREILKEPVHGVYY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0202)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (308-317)

**Author Location RT (LAI)** 

Epitope EILKEPVGHV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**References** Menendez-Arias *et al.* 1998; van der Burg *et al.* 1997

- Recognized by CTL from a long-term survivor, SPIETVPVKL was also recognized.
- Recognized by CTL from a progressor, EELRQHLLRW and TWETWWTEYW were also recognized.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A\*02)

Keywords HAART, ART

References Huang et al. 2000

- The single cell ELISPOT assay was optimized and highly specific, and found to work well even after the primary cells had been frozen and thawed.
- Increases in gamma interferon producing cells were observed in response to anti-retroviral therapy using single cell IFNgamma-production ELISPOT.

**HXB2 Location** RT (309–317)

Author Location RT (476-484)

**Epitope** ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A\*02)

**Keywords** HAART, ART

References Rinaldo et al. 2000

Administration of triple-drug antiretroviral therapy (IDV, 3TC and ZDV) sometimes showed a transient increase and other times failed to increase CTL responses in patients with advanced HIV disease, but there is a stable population of tetramer stained HIV-specific CD8+ CD45RO+ cells that persist after therapy and long periods of virus being below the level of detection.

**HXB2 Location** RT (309–317)

**Author Location RT** 

Epitope ILKEPVHGV

Epitope name IV9

Immunogen HIV-1 infection

Species (MHC) human (A\*02)

Keywords HAART, ART, immunodominance

References Scott-Algara et al. 2001

- This study examined with CTL response in HLA A\*02+ children by tetramer staining for HLA-A2 immunodominant epitopes SLYNTVATL and ILKEPVHGV.
- 71% of the 28 HIV-1 infected HLA-A\*02 positive children recognized both epitopes, with cells from 26 children stained positive by the gag tetramer (SLYNTVATL) and 21 children by the pol tetramer (ILKEPVHGV)
- There were no differences observed in children that had therapy versus those that did not.
- Tetramer-binding cells were memory activated CD28-, CD45RO+, CD45RA- HLADR+, CD69-, CD8+ T-cells.

**HXB2 Location** RT (309–317)

**Author Location RT** 

Epitope ILKEPVHGV

Epitope name POL

Immunogen HIV-1 infection

Species (MHC) human (A\*02)

Country France.

Assay type Cytokine production, Tetramer binding, In-

tracellular cytokine staining, Flow cytometric T-cell cytokine assay

Keywords responses in children, characterizing CD8+ T

cells

References Scott-Algara et al. 2005

 Only a fraction of HIV-1-specific CD8 T-cells detected in the PBMC of 17 infected children (ages 2-18) were able to produce cytokines (IFN-gamma, TNF-alpha) or chemokines (CCL4, CCL5) after stimulation with the cognate peptide. A negative correlation was found between the plasma viral load and the precentage of CD8+ Gag-specific T-cells secreting IFN-gamma. Tetramers used in this study were SLYNTVATL-HLA-A\*02 and ILKEPVHGV-HLA-A\*02.

**HXB2 Location** RT (309–317)

**Author Location** 

Epitope ILKEPVHGV Immunogen HIV-1 infection Species (MHC) human (A\*0201) Keywords acute/early infection References Wilson *et al.* 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and a close temporal relationship between the number of circulating HIV-specific T cells and viral load.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.

- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPOEVVL.

**HXB2 Location** RT (309–317)

Author Location Pol (476–484)

**Epitope** ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Spiegel et al. 2000

- High levels of CD8+ HIV-1 specific and cytomegalovirus specific CTL were detected by HLA-A\*0201-peptide tetramers in 3 infected subjects with very low CD4 counts, but CD8 T cell mediated effector activity was not seen.
- Thus HIV-1 specific CD8+ cells may be present but may lack direct effector activity in late disease, suggesting that overcoming antigen unresponsiveness may be a useful therapeutic strategy.

**HXB2 Location** RT (309–317)

**Author Location** Pol (476–484)

**Epitope** ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** epitope processing, immunodominance

References Sewell et al. 1999

- Proteasome regulation influences epitope processing and could influence immunodominance.
- The proteasome is inhibited by lactacystin treatment, and gamma IFN induces expression of proteasome subunits, LMP2 and LMP7, which combine with the proteasome to create an immunoproteasome.
- IFN-gamma induction of the immunoproteasome and lactacystin inhibition increases the presentation of the A\*0201 VIYQYMDDL epitope, but decreases the presentation of the A\*0201 ILKEPVHGV epitope, which is immunodominant within pol proteins, showing the two epitopes are processed by different pathways.
- ILKEPVHGV seems to be processed by the classical proteasome pathway, while VIYQYMDDL appears to be destroyed by this pathway.
- This epitope contains the catalytic site (YMDD) of RT, a conserved sequence in HIV-1 which restricts escape mutants.

**HXB2 Location** RT (309–317)

**Author Location** Pol (476–484)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** epitope processing

References Loing et al. 2000

The ILKEPVHGV was modified by the addition of an N-palmitoyl-lysine residue at the P0, P1 or P10 positions of the parent peptide to create a lipopeptide for direct antigen delivery to the cytoplasm for processing.

tional CTL recognition up to 48 hours in comparison to the parent peptide.

**HXB2 Location** RT (309-317) Author Location Pol (510-518) Epitope ILKEPVHGV Immunogen vaccine

Vector/Type: canarypox, vaccinia HIV com-

ponent: Env, Gag, Nef, Pol

Species (MHC) human (A\*0201) References Larsson et al. 1999

• ELISPOT was used to assay the CD8 T cell response to the HIV-1 proteins Gag, Pol, Nef or Env expressed in vaccinia or canarypox vectors in 19 HIV+ people.

- The highest CTL frequency was directed at epitopes in Pol.
- In A\*0201 individuals, higher numbers of spot-forming T cells were directed against HIV-1 proteins expressed in vaccinia than to peptides SLYNTVATL and ILKEPVHGV presented by A2.

**HXB2 Location** RT (309–317) Author Location RT (476–484) Epitope ILKEPVHGV Immunogen HIV-1 infection Species (MHC) human (A\*0201) Keywords TCR usage References Wilson et al. 1998a

- HIV+ individuals were followed longitudinally using MHC tetramers in combination with 14 anti-BV chain MAbs, and clonal expansion of HIV-specific T cells was followed in vivo.
- Seven HIV+ people were studied, and all showed expansions of particular TCR BV clones, often several, relative to uninfected controls.
- Three patients were followed in detail, TCR VB expansions persisted for 2 to 3 years, with occasional transient increases.

**HXB2 Location** RT (309-317) **Author Location** RT (476–484) Epitope ILKEPVHGV Immunogen HIV-1 infection Species (MHC) human (A\*0201) Keywords immunodominance References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 2/11 of the A2+ individuals responded to ILKEPVHGV, and neither of these two responded to SLYNTVATL.

**HXB2 Location** RT (309–317) Author Location Pol **Epitope ILKEPVHGV** Immunogen HIV-1 infection Species (MHC) human (A\*0201) Keywords HAART, ART References Gray et al. 1999

• The N-terminal modification increased the life span for func• Administration of highly active antiretroviral therapy (HAART) reduced CD8+ cell frequency, and the CD8+ cells detected by tetramer staining were likely to be memory cells, indicating that persistently replicating viral populations are needed to maintain high frequencies of HIV-1 specific CTL.

> HXB2 Location RT (309-317) Author Location RT (476-484) **Epitope ILKEPVHGV** Immunogen HIV-1 infection Species (MHC) human (A\*0201)

> > References Menendez-Arias et al. 1998; Ogg et al. 1998b

- HLA-tetrameric complexes were used in a cross-sectional study of 14 untreated HLA A\*0201 positive individuals, revealing an inverse relationship between HIV Gag and Pol specific CTL effector cells (CTLe) and viral load.
- Inclusion of both the p17 SLYNTVATL and RT ILKEPVHGV epitopes gives a good representation of HLA A\*0201-restricted activity.
- No correlation was observed between the CTLe and CD4 count or clearance rate of productively infected cells.

HXB2 Location RT (309-317) **Author Location RT Epitope** ILKEPVHGV Immunogen vaccine Vector/Type: vaccinia

Species (MHC) human (A\*0201)

References Hanke et al. 1998a; Hanke et al. 1998b

 This epitope was shown to be processed and presented to appropriate CTL clones upon infection of human target cells with vaccinia virus Ankara (VVA) carrying 20 HIV-1 epitopes recognized by humans.

**HXB2 Location** RT (309–317) Author Location RT (476-484) Epitope ILKEPVHGV Immunogen in vitro stimulation or selection Species (MHC) human (A\*0201) Keywords binding affinity

References Konya et al. 1997; Menendez-Arias et al.

- This epitope was included as a positive control.
- Binding affinity to A\*0201 was measured,  $C_1/2\max \mu M = 12$

**HXB2 Location** RT (309–317) **Author Location** RT (468–476) Epitope ILKEPVHGV

**Immunogen** in vitro stimulation or selection

Species (MHC) human (A\*0201)

References van der Burg et al. 1996

- · Immunogenic in humans, slow dissociation rate, and associated with immunogenicity in transgenic HLA-A\*0201/K<sup>b</sup> mice.
- CTL generated by in vitro stimulation of PBMC derived from uninfected individual.

**HXB2 Location** RT (309-317) **Author Location** RT (468–476) **Epitope ILKEPVHGV** 

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

References van der Burg et al. 1995

• Binds HLA-A\*0201 – CTL generated by *in vitro* stimulation of PBMC from an HIV negative donor.

**HXB2 Location** RT (309-317)

Author Location RT (476-484)

**Epitope** ILKEPVHGV **Immunogen** HIV-1 infection

Species (MHC) human (A\*0201)

References Menendez-Arias et al. 1998; Pogue et al. 1995

 Mutational study: position 1 I to Y increases complex stability with HLA-A\*0201.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484)

**Epitope** ILKEPVHGV **Immunogen** HIV-1 infection

**Species** (MHC) human (A\*0201) **Keywords** review, escape

**References** Goulder *et al.* 1997e; Goulder *et al.* 1997a; Menendez-Arias *et al.* 1998

- HLA-identical sibling hemophiliac brothers were both infected with the same batch of factor VIII.
- One had a response to gag A2 epitope SLYNTVATL, the other to pol A2 epitope ILKEPVHGV. They were tested 6-8 years after infection.
- Viral sequencing from the twin that had no response to SLYNT-VATL indicated his virus had the substituted form SLH-NAVAVL.
- 71% of an additional set of 22 HIV-1 infected HLA-A\*0201 positive donors preferentially responded to gag SLYNTVATL.
- Those individuals with a pol ILKEPVHGV response tended to have mutations in or around SLYNTVATL.
- Goulder *et al.* [1997a] is a review of immune escape that summarizes this study.

   A minigene vaccine construct encoding 6 HLA 2.1 and 3 HLA All restricted CTL epitopes, the universal Th cell epitope

**HXB2 Location** RT (309–317)

Author Location RT (309-317)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

References Altman et al. 1996

- This paper introduces the tetramer methodology which permits quantification of specific CTL based on expression of specific TCRs—HLA-A2 tetramers were prepared that can stain CTL lines specific for ILKEPVHGV and SLYNTVATL, and can quantify HIV-specific CD8+ cell lines in freshly isolated PBMCs.
- Three patients only stained the Gag epitope SLYNTVATL, one patient had the highest frequency of tetramer staining to the Pol epitope (0.77%), less to the Gag epitope (0.28%)
- The A2-Pol CD8+ clones were CD45RO positive and HLA-DR and CD38 negative, suggesting a memory rather than effector phenotype.

**HXB2 Location** RT (309–317)

Author Location RT (476-484)

**Epitope** ILKEPVHGV

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Keywords epitope processing

**References** Menendez-Arias *et al.* 1998; Walter *et al.* 1997

- HLA-A2 heavy chain and β2-microglobulin expressed in E. coli were refolded in the presence of this peptide.
- The HLA-A2-peptide complex elicited HLA-A2 peptidespecific CTL response in cells lacking HLA-A2.
- Suggests that preformed HLA-peptide complexes could provide an alternate to intracellular processing for immunogens.

**HXB2 Location** RT (309–317)

Author Location RT (464-472)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords HAART, ART

References Gray et al. 1999

- Peptide-tetramer complexes of A\*0201 and SLYNTVATL or ILKEPVHGV were used to study individuals receiving HAART to determine the frequency of Class I HLA-restricted anti-HIV CD8+ T cells.
- 17/18 asymptomatic patients had a CTL response to one or both epitopes 72% had a CTL response to SLYNTVATL.
- After HAART, the majority of the epitope-specific CTL were apparently memory cells.

**HXB2 Location** RT (309–317)

**Author Location** Pol

Epitope ILKEPVHGV

Immunogen vaccine

Vector/Type: DNA

Species (MHC) transgenic mouse (A\*0201)

References Ishioka et al. 1999

- A minigene vaccine construct encoding 6 HLA 2.1 and 3 HLA A11 restricted CTL epitopes, the universal Th cell epitope PADRE (pan-DR epitope) and an ER translocating signal sequence was constructed.
- The epitopes were chosen for dominant recognition by CTLs during HBV and HIV infections in humans.
- HLA transgenic mice were used for quantitating in vivo immunogenicity of DNA vaccines encoding HLA-restricted CTL epitopes strong responses were observed to all nine epitopes, and CTL memory persisted up to four months after a single injection.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords escape

References Brander et al. 1998a

- Of 17 infected HLA A\*0201 subjects, 13 had CTL responses against the p17 SLYNTVATL epitope, six recognized ILKEPVHGV and five recognized VIYQYMDDL, and there was no correlation between viral load and recognition of a specific epitope or evidence of immune escape.
- Only one subject had CTL against all three epitopes.

- Subjects were part of the San Francisco City Clinic Cohort, the ARIEL project and from the Boston area.
- C. Brander notes this is an A\*0201 epitope.

HXB2 Location RT (309–317)
Author Location Pol (476–484)
Epitope ILKEPVHGV
Immunogen HIV-1 infection
Species (MHC) human (A\*0201)
Keywords HAART, ART
References Ogg et al. 1999

- CTL effector levels were measured after potent ARV therapy using HLA-tetramer complexes for the A\*0201 epitopes SYLVTVATL and ILKEPVHGV in seven patients, and the B\*3501 epitope DPNPQEVVL in one additional patient.
- Levels of CTL effectors typically decline for 5-7 days and then rebound, fluctuating during the first two weeks of therapy.
- After the early fluctuation, there was a steady exponential decay with a median half-life of 45 days.

HXB2 Location RT (309–317) Author Location RT (476–484 LAI) Epitope ILKEPVHGV Subtype B

Immunogen HIV-1 infection
Species (MHC) human (A\*0201)
Keywords optimal epitope
References Frahm et al. 2007

• C. Brander notes this is a A\*0201 epitope.

HXB2 Location RT (309–317) Author Location RT (476–484) Epitope ILKEPVHGV

Epitope name IV9

**Immunogen** HIV-1 infection, in vitro stimulation or selection

**Species (MHC)** human (A\*0201) **References** Dela Cruz *et al.* 2000

- Integration of HIV RT CTL epitopes into the N-terminus of the HLA-A2 heavy chain, or tethering the epitopes to the target chain, resulted in epitope-specific lysis by CD8+ CTL.
- These antigens could also be used to stimulate primary responses *in vitro*.

HXB2 Location RT (309–317) Author Location RT (309–317) Epitope ILKEPVHGV Epitope name P1

Immunogen HIV-1 infection Species (MHC) human (A\*0201) Keywords HAART, ART, escape References Samri et al. 2000

• The epitope was recognized by patient 250#0 but not in another A\*0201+ patient, 201#5, in a study of the effects of therapy escape mutations on CTL recognition.

HXB2 Location RT (309–317) Author Location Pol (LAI) Epitope ILKEPVHGV Subtype B Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Keywords dendritic cells

References Engelmayer et al. 2001

- Recombinant canarypox virus vector containing HIV-1 sequences, upon infection of mature dendritic cells, can trigger specific lysis through in vitro by T-cells from HIV-1 infected individuals at levels comparable to the response seen to HIV carried in vaccinia vectors.
- Recombinant canarypox virus vector containing HIV-1 sequences can also stimulate HIV-specific CD4+ helper T-cell responses.

HXB2 Location RT (309–317)

**Author Location** Pol

Epitope ILKEPVHGV Immunogen HIV-1 infection Species (MHC) human (A\*0201)

References Gea-Banacloche et al. 2000

- In a study including many long-term non-progressors, no correlation between plasma virus levels and number of HIV-specific CD8+ T-cells was found.
- High frequencies of circulating CD8+ T-cells were HIV-1 specific, and the majority of these responses were to gag-pol gene products.
- 4/21 subjects were HLA-(A\*0201), and of these only 2 subjects (patient 3 and 19) tested positive to this epitope.

HXB2 Location RT (309–317) Author Location Pol (476–484) Epitope ILKEPVHGV

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Keywords HAART, ART, rate of progression

References Jin et al. 2000a

- The CTL effector levels (CTLe) were compared in long term non-progressors (LTNP) with low viral load and in patients whose virus was well-suppressed by therapy, using a tetramer assay.
- LTNPs have high memory CTLe numbers and low viral load, while HAART patients had low CTLe numbers and low viral load.

**HXB2 Location** RT (309–317) **Author Location** Pol (476–484)

Epitope ILKEPVHGV

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

References Appay et al. 2000

- Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virus-specific CD8+ T cells produced IFN- $\gamma$  and MIP-1 $\beta$  with a distinct subset that failed to produce TNF- $\alpha$

**HXB2 Location** RT (309–317)

Author Location Pol

**Epitope** ILKEPVHGV **Immunogen** HIV-1 infection

Species (MHC) human (A\*0201)

Keywords dendritic cells

References Ostrowski et al. 2000

- The role of CD4+ T-cell help in expansion of virus-specific memory CTL was studied through co-culture ex vivo
- Optimal expansion of HIV-1-specific memory CTL depended on CD4+ T cell help in 9 of 10 patients CD40 ligand trimer (CD40LT) could enhance CTL in the absence of CD4+ T cell help to a variable degree in most of patients.
- Those CTL that didn't respond to CD40LT could expand with IL2 present, and IL15 produced by dendritic cells also contributes.
- The T-helper epitope used for CD4+ T cell stimulation was the universal tetanus helper epitope TET830-843 (QYIKANSK-FIGITE)

**HXB2 Location** RT (309–317)

Author Location RT (309-317)

**Epitope** ILKEPVHGV

**Epitope name** RT2

Immunogen vaccine

Vector/Type: HIV-1 peptide in filamentous bacteriophage major coat protein HIV component: RT

 $\textbf{Species} \ (\textbf{MHC}) \ \ \text{human, transgenic mouse} \ (A*0201)$ 

References Guardiola et al. 2001

 HLA-A2 transgenic mice were injected with bacteriophage antigens expressing a Th epitope and the HIV CTL epitope ILKEPVHGV, and epitope-specific cytotoxic activity was induced.

**HXB2 Location** RT (309–317)

**Author Location** Pol (476–484)

**Epitope** ILKEPVHGV

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

**Keywords** epitope processing, immunodominance

References Sewell et al. 2002

- Epitope processing of three different HLA-A\*0201 HIV epitopes was shown to use different pathways, which might influence patterns of immunodominance. .174 cells were used that lack TAP1 and TAP2 genes, as well as the LMP2 and LMP7 genes that encode the beta-subunits of the immunoproteasome. These genes could be added back through transfection to study processing.
- ILKEPVHGV was efficiently presented in TAP-1 and -2 transfected cells while VIYQYMDDL and SLYNTVATL were not. VIYQYMDDL was destroyed by the MB1 subunit of the protease, and could be expressed in the presence of the proteasome inhibitor lactacystin, but SLYNTVATL expression was not restored. SLYNTVATL expression was unaltered by lactacystin in a wild type cell line.

**HXB2 Location** RT (309–317)

Author Location Pol

**Epitope** ILKEPVHGV

**Epitope name** IL-9

Immunogen HIV-1 infected monocyte-derived

Species (MHC) mouse (A\*0201)

References Poluektova et al. 2002

- Nonobese diabetic NOD-C.B-17 SCID mice were reconstituted with HLA-A\*0201 positive human PBL and injected with HIV-1 infected monocyte-derived macrophages MDM in the basal ganglia to provide a mouse model of HIV-1 encephalitis.
- HLA-A\*0201 CTL responses were detected by tetramer staining in the spleen in seven days, increased through day 14, and the numbers of productively infected were reduced >85% in the second week.

**HXB2 Location** RT (309–317)

Author Location RT (476–484 LAI)

Epitope ILKEPVHGV

Epitope name LR22

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant (IFA), Montanide (ISA 720), P30, PLG

Species (MHC) mouse (A\*0201)

References Peter et al. 2001

**Keywords** binding affinity, vaccine-specific epitope characteristics improved deminance

acteristics, immunodominance

- The stability of peptide binding to HLA-A2.1 was determined for six HLA-A2.1 peptides included in this vaccine study
   ILKEPVHGV (RT), SLYNTVATL (p17), SLLNATDIAV (gp41) and LLWKGEGAV (RT) all bound with high affinity comparable to a influenza epitope reference (GILGFVFTL), while RGPGRAFVTI and VIYQYMDDL bound with a lower affinity (relative binding activity = 0.01).
- The four high-affinity peptides formed stable complexes with half-lives ranging between 8 and 32 hours, while the low affinity peptides had half lives of less than an hour.
- HLA-A2.1 transgenic mice were immunized with the six HIV-1 peptides and P30, as a universal T-helper epitope, with IFA or Montanide or microspheres as adjuvants.
- All peptides except VIYQYMDDL induced a stong CTL response in Cr-release assays stronger responses were observed when peptides were delivered alone, indicating immunodominance when the combination was used.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484 LAI)

Epitope ILKEPVHGV

Epitope name LR22

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant

(IFA), IL-12, P30

Species (MHC) mouse (A\*0201)

Keywords vaccine-specific epitope characteristics, im-

munodominance

References Peter et al. 2002

• When HIV-1 peptides were used to vaccinate HLA-A2.1 transgenic A2-Kb mice, strong responses to five peptides were observed when the peptides were given individually, but immunodominance limited the response to some of the peptides when they were given in combination Peter *et al.* [2001]. IL-12 can counteract immunodominance in BALB/c mice, so it was given with the multiple epitope vaccination, and was instead found to specifically eliminate the HLA-A2.1-epitope CTL responses, but not Kb CTL responses. This was possibly a consequence of transient depletion of T-cells, B cells and macropahges in the spleen.

HXB2 Location RT (309–317) Author Location RT (309–317)

**Epitope** ILKEPVHGV

Subtype B

Immunogen vaccine

Vector/Type: peptide HIV component: RT Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) transgenic mouse (A\*0201)

**Keywords** binding affinity, vaccine-specific epitope characteristics

References Boissonnas et al. 2002

 Ten naturally occurring variants of the Nef epitope VLMWQFDSRL were tested for their affinity to HLA-A\*0201 and for their ability to induce gamma-IFN and cytotoxic functions through vaccination of HLA-A\*0201 transgenic mice.

 ILKEPVHGV could induce HLA-A\*0201 vaccine responses, and was a positive control.

**HXB2 Location** RT (309–317)

**Author Location** Pol (468–476)

Epitope ILKEPVHGV

Immunogen vaccine

Vector/Type: DNA HIV component: HIV-1

Species (MHC) mouse (A\*0201)

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance

References Singh et al. 2002; Sykes & Johnston 1999

- C3H (H-2k) transgenic mice carrying a fused HLA-A\*0201 alpha1 and alpha2 and H-2Dk alpha3 hybrid class I molecule were immunized using an epidermal gene gun with an ubiquitin expression library of 32 plasmids that spanned the HIV-1 genome. Ubiquitin targets the expressed HIV-1 peptides to the proteasome.
- A single immunization with the UB-HIV-1 library vaccine induced potent, stable and multivalent CTL responses against all library members.
- Immunodominant epitopes SLYNTVATL (Gag), ILKEPVHGV (Pol), RIQRGPGRAFVTIGK (Env) and AFHHVAREK (Nef) elicited strong CD8+/IFN- responses and stimulated CTL that were functional in a Cr-release assay and against wild type antigen.
- The presence of multiple plasmids HLA-A\*0201-restricted CTL epitopes did not decrease CTL immunogenicity, and CTL responses to single peptide immunizations were comparable to responses based on mixtures of either 16 or 32 peptides.

**HXB2 Location** RT (309–317)

**Author Location** Pol

Epitope ILKEPVHGV

Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

**Species (MHC)** human, macaque (A\*0201)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

**HXB2 Location** RT (309-317)

Author Location Pol (476-484)

**Epitope ILKEPVHGV** 

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Keywords epitope processing, dendritic cells

References Andrieu et al. 2003

- This study demonstrates that lipopeptides carrying epitopes can be taken up by human dendritic cells, processed using different pathways, and recognized by epitope-specific CD8+T-cells originally derived from HIV+ individuals. The RT ILKEPVHGV peptide was embedded in a longer peptide fragment in the lipopeptide, and was internalized by endocytosis and processed in the cytosol by proteasomal cleavage by following an endosome-to-cytosol pathway for processing and presentation. Administration of epoxomycin, a proteasome inhibitor, completely abrogated epitope presentation to a CD8+T-cell line, while monensin, an inhibitor of acid-dependent endosomal enzyme activity did not.
- In contrast to the RT epitope, dendritic cell presentation of the Nef epitope QVPLRPMTYK embedded in a longer peptide in a lipopeptide was not inhibited by epoxomycin, but was inhibited by monensin, indicative of endocytolitic epitope processing.

**HXB2 Location** RT (309–317)

**Author Location** 

**Epitope** ILKEPVHGV

Epitope name IV9 Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Assay type Cytokine production, proliferation, Tetramer binding, Intracellular cytokine staining,

Chromium-release assay

References Dagarag et al. 2003

- Telomer length is short in the CD8+ T-cell compartment of HIV-1 infected people, indicating excessive CTL activation and premature senescense. Here human telomerase RT (hTERT) transduction of HIV-1-specific CTL was used to study the functional impact of telomerase. Telomerase expression enhanced proliferative capacity, as well as cytolytic and antiviral capabilities; cytokine production was unchanged. hTERT transduced CTLs were 10-fold more efficient in controlling HIV-1 replication in culture. Thus telomerase transduction can restore CTL mediated cytolysis, and may have therapeutic potential.
- Three polyclonal CD8+ T-cell lines derived from an HIV-1, HLA A\*0201 positive patient were used in this study, including one specific for this epitope. An IV9-specific monoclonal cell line, 68A62 was also generated.

**HXB2 Location** RT (309–317)

**Author Location** Pol (464–472)

**Epitope** ILKEPVHGV

Epitope name 19V Subtype B Immunogen vaccine

*Vector/Type:* peptide *HIV component:* RT *Adjuvant:* CpG immunostimulatory sequence (ISS)

Species (MHC) transgenic mouse (A\*0201)

Donor MHC H-A2/Kb

Assay type Cytokine production, Tetramer binding, Intracellular cytokine staining, Chromium-release assay

References Daftarian et al. 2003

- HLA-A\*0201 transgenic mice were immunized with a Th-CTLfusion peptide composed of the I9V CTL epitope linked to the promiscuous PADRE Th epitope. The peptide only when given in combination with CpG elicited strong I9V-CTL responses.
- The peptide-CpG vaccinated mice, when challenged with pol embedded in vaccinia (pol-vv), could clear the virus from the ovaries. Additionally, intranasal immunized mice given an intranasal pol-vv challenge reduced virus in the lungs.

**HXB2 Location** RT (309–317)

**Author Location** Pol (476–484)

Epitope ILKEPVHGV

Epitope name IV9

Immunogen HIV-1 infection Species (MHC) human (A\*0201) Assay type Tetramer binding

Keywords genital and mucosal immunity

References Shacklett et al. 2003

- Lymphocytes from rectal biopsies were used to characterize the CD8+T cell response to HIV in GALT, Gut-associated lymphoid tissues. Patients were selected on the basis of being HLA-A2+ and having detectable SLYNTVATL and ILKEPVHGV tetramer responses in PBMC. SLYNTVATL frequency was increased in GALT relative to PBMC in 6/7 patients studied, while a control response to a CMV-peptide was diminished in GALT. Only two patients had ILKEPVHGV CD8+ T cell responses, and both had slightly higher frequencies in GALT than PBMC.
- HIV may perturb lymphocyte retention in GALT, suggested by an overall reduction of GALT CD8+ cells expressing alphaEbeta7. GALT HIV-specific CD8+ T cells expressed alphaEbeta7, suggesting mucosal priming.

**HXB2 Location** RT (309–317)

**Author Location** RT (309–317 MN)

Epitope ILKEPVHGV

Subtype B

Immunogen vaccine

Vector/Type: DNA, polyepitope Strain: B clade MN HIV component: gp120, Protease, RT Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) humanized mouse (A\*0201)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance, immunotherapy

References Isaguliants et al. 2004

- Immunization of HLA-A\*0201-transgenic mice with synthetic genes encoding clusters of human A\*0201 CTL epitopes located at the sites of drug resistance mutations, induced RTspecific cellular responses indicating the immunogenicity of these constructs. This vaccine strategy may be a first step towards a therapeutic vaccine against drug-resistant strains.
- This was one of five HLA-A\*0201 epitopes from the RT or
  protease proteins that was included in the polyepitope vaccine.
  When the transgenic HLA A\*0202 mice were vaccinated with
  the polyepitope construct or with a mixture of RT peptides,
  a sustained low level CD8+ T-cell gamma IFN response was
  observed, in contrast to when an intact RT gene was used for
  vaccination.

**HXB2 Location** RT (309-317)

**Author Location** 

Epitope ILKEPVHGV

**Epitope name** IV9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, T-cell Elispot,

Flow cytometric T-cell cytokine assay

**Keywords** epitope processing, escape, kinetics, variant

cross-recognition or cross-neutralization

References Jamieson et al. 2003

 Epitope escape mutations in chronically infected individuals developed over several years indicating slight selective advantage of escape mutants. The maturation state of CTLs appear to affect the rate of epitope mutation and CTL decay. • In two patients, IV9 mutations preceded the loss of IV9-specific CD8+ T-cells. In a third patient, escape mutations were coincident with IV9-specific CD8+ T-cell loss. One patient was infected with a ilepvhgA variant, and transiently reverted to the consensus form at year 3. One patient never made a response to IV9 despite being infected with the consensus form of the epitope.

**HXB2 Location** RT (309-317)

**Author Location** Gag

**Epitope** ILKEPVHGV

Epitope name IV9

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Assay type Tetramer binding, Chromium-release assay,

Flow cytometric T-cell cytokine assay

Keywords epitope processing, rate of progression, immunodominance, acute/early infection, den-

dritic cells, TCR usage, memory cells

References Kan-Mitchell et al. 2004

• In contrast to IV9-CTLs, SL9-CTLs were shown to be primed by immature DCs and to be independent of help from CD4+ or exogenous IL2 and sensitive to paracrine IL-2-induced apoptosis.

**HXB2 Location** RT (309–317)

Author Location Pol (468–476 IIIB)

Epitope ILKEPVHGV

Epitope name pol468-476

Subtype B

Immunogen vaccine

HIV component: Gag-Pol

**Species (MHC)** humanized mouse (A\*0201)

Assay type Intracellular cytokine staining

**Keywords** epitope processing, vaccine-specific epitope

characteristics, variant cross-recognition or cross-neutralization, vaccine antigen design

References Singh & Barry 2004

- When A\*0201-C3H/J transgenic mice were immunized with plasmids encoding wild-type gag-pol, codon-optimized (CO) gag-pol, and an expression library vaccine expressing 16 fragments of gag-pol fused with ubiquitin for proteosome targeting (ELI), the ELI vaccine produced up to 10-fold higher CD8 T-cell responses than the other two vaccines. In contrast to the wt and CO vaccines, which tended to augment only immunodominant responses, boosting with the ELI vaccine resulted in many CD8 responses against variant epitopes from different HIV-1 clades, and against drug-resistant variants.
- · This epitope was recognized in transgenic mice vaccinated with all three vaccine constructs, but the most intense responses were to the ELI vaccine.

**HXB2 Location** RT (309–317)

Author Location Pol (464–472)

Epitope ILKEPVHGV

Epitope name IV9

Subtype B

Immunogen HIV-1 infection

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Species (MHC) human (A*0201)
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**Donor MHC** A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, Chromium-release assay

**Keywords** escape, acute/early infection

References Bansal et al. 2005

- Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- The response to this epitope was not detected until month 25 and increased over time.

**HXB2 Location** RT (309-317)

**Author Location** Pol

Epitope ILKEPVHGV

Epitope name 19V

Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV component: gp140, gp140ΔV3

**Species (MHC)** transgenic mouse (A\*0201)

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay

Keywords memory cells, vaccine antigen design, anti-

body generation, characterizing CD8+ T cells

References Lorin et al. 2005

Vector/Type: DNA Strain: B clade IIIB • A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted CTL responses against multiple conserved HIV-1 epitopes in HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw virus expressing gp140 with deleted V1, V2, and V3 loops successfully induces neutralizing antibodies against HIV-1. A live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination against HIV and measles.

**HXB2 Location** RT (309–317)

Author Location RT (476-484 LAI)

Epitope ILKEPVHGV

Epitope name P1

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (A\*0201, A\*0205)

Keywords HAART, ART

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using tetramer staining or CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened - eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+

PBL – but with continued viral suppression, HIV-specific responses diminished.

 Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

**HXB2 Location** RT (309–317)

**Author Location** Pol (476–484)

Epitope ILKEPVHGV

Immunogen vaccine

Vector/Type: vaccinia

Species (MHC) human (A2)

References Woodberry et al. 1999

- A polyepitope vaccine was generated in a vaccinia construct that contiguously encoded seven epitopes, all presented by HLA A-2.
- HHD mice have a transgene of HLA A2 linked to the transmembrane and cytotoxic domains of H-2D<sup>d</sup> – this transgene is the only MHC molecule expressed in the mice.
- CTL responses to Gag (77-85) SLYNTVATL, Pol (476-484) ILKEPVHGV, gp120 (120-128) KLTPLCVTL, and Nef (190-198) AFHHVAREL were observed in HIV polytope HHD-vaccinated mice, and these responses were enhanced with vaccinia boost.
- No CTL immune responses were generated against HLA A2restricted HIV epitopes Nef 157-166 (PLTFGWCYKL), Pol 346-354 (VIYQYMDDL), and Nef 180-189 (VLEWRFDSRL)
- Sixteen HLA A2+ patients were tested for their ability to make CTL responses by peptide restimulation in culture with the epitopes selected for inclusion in the polytope – one individual recognized all seven of these epitopes; 7 patients had CTL cultures able to recognize at least one of the epitopes, and 6 of those 7 recognized more than one epitope, but they were not able to test all peptides for all patients; many patients only had three peptides tested.
- ILKEPVHGV was recognized by 2 of the patients.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484)

Epitope ILKEPVHGV

Immunogen HIV-1 infection Species (MHC) human (A2)

Keywords subtype comparisons, TCR usage

References Kolowos et al. 1999

- TCR usage in CTL specific for this epitope was examined in three patients and identical V $\beta$ 6.1 and Valpha2.5 gene segments were used and two of the patients had very similar complementarity-determining regions clonal expansion of RT-HIV-specific CTL can contribute to the skewed TCR repertoire in HIV-1 infected patients.
- CTL clones from all three patients showed similar sensitivity to mutation in the epitope, ilkepvhEv was well recognized (the sequence from SF2), ilkDpvhgv was not (the common A clade form)

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484)

**Epitope** ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Collins et al. 1998

- Nef down-regulates MHC class I molecules, which inhibits CTL killing of HIV-infected targets.
- The anti-RT CTL clone killed Nef- cells less efficiently than anti-gag clones, correlated with the reduced expression of RT.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484 LAI)

**Epitope** ILKEPVHGV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords dendritic cells

References Fan et al. 1997

 The capacity of dendritic cells to process and present antigen and stimulate anti-HIV-1 CTL memory responses was studied.

**HXB2 Location** RT (309–317)

**Author Location** RT (464–472)

**Epitope** ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords dendritic cells

References Kundu et al. 1998b

- Allogeneic dendritic cells (DCs) were obtained from HLAidentical siblings, pulsed with rgp160 MN or A2-restricted HIV-1 epitope peptides, and infused monthly into six HIVinfected patients.
- 1/6 showed increased env-specific CTL and increased lymphoproliferative responses, 2/6 showed increase only in proliferative responses, and 3/6 showed no change – pulsed DCs were well tolerated.
- ILKEPVHGV is a conserved HLA-A2 epitope included in this study 5/6 patients had this sequence as their HIV direct sequence, and these had a detectable CTL response—one person carried the form ILREPVHGV and had no detectable CTL.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484)

**Epitope** ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

**References** Menendez-Arias *et al.* 1998; Tsomides *et al.* 

 CTL clones recognize naturally processed peptide – peptide abundance corresponded to level of CTL killing.

**HXB2 Location** RT (309–317)

Author Location RT (476–484)

Epitope ILKEPVHGV

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2)

**Keywords** subtype comparisons, HIV exposed persistently seronegative (HEPS)

**References** Rowland-Jones *et al.* 1998a

A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades – such cross-reactivity could protect against both A and D and confer pro-

tection in Nairobi where both subtypes are circulating.

- The A subtype consensus is ILKDPVHGV.
- The D subtype consensus is identical to the epitope ILKEPVHGV.

**HXB2 Location** RT (309-317)

Author Location RT (476–484)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** subtype comparisons

References Cao et al. 1997a; Menendez-Arias et al. 1998

- The consensus peptides of B and D clade viruses and some As have the sequence ILKEPVHGV.
- The consensus peptide of a subset of A clade viruses, ILKD- Precise identification of the nonamer that binds to A2. PVHGV, is not cross-reactive.

**HXB2 Location** RT (309–317)

Author Location RT (476–484)

**Epitope ILKEPVHGV** 

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Menendez-Arias et al. 1998; Yang et al. 1996

- CD4+ cell lines acutely infected with HIV were studied to determine their susceptibility to lysis by CTL.
- Clones specific for RT lysed HIV-1 infected cells at lower levels than Env or Gag specific clones.
- The distinction was thought to be due to lower expression of RT relative to Env and Gag.
- CTL can lyse infected cells early after infection, possibly prior to viral production.

**HXB2 Location** RT (309-317)

Author Location RT (476-484)

**Epitope ILKEPVHGV** 

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type CTL suppression of replication

References Yang et al. 1997a

- CTL inhibit HIV-1 replication at effector cell concentrations comparable to those found in vivo.
- CTL produced HIV-1-suppressive soluble factors MIP-1 $\alpha$ , MIP-1 $\beta$ , RANTES, after antigen-specific activation.
- CTL suppress HIV replication more efficiently in HLAmatched cells.

**HXB2 Location** RT (309–317)

**Author Location** RT (309–317)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords TCR usage

References Menendez-Arias et al. 1998; Moss et al. 1995

 Two clones were obtained with different TCR usage, V<sub>B</sub>1 and  $V_{\beta}21.$ 

**HXB2 Location** RT (309–317)

Author Location RT (476–484)

**Epitope ILKEPVHGV** 

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Menendez-Arias et al. 1998; Musey et al.

· Cervical CTL clones from an HIV-infected woman recognized this epitope.

**HXB2 Location** RT (309-317)

Author Location RT (476–484 LAI)

Epitope ILKEPVHGV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Menendez-Arias et al. 1998; Tsomides et al. 1991

**HXB2 Location** RT (309–317)

Author Location RT (476–484 LAI)

**Epitope** ILKEPVHGV

Subtype B

Immunogen peptide-HLA interaction

Species (MHC) human (A2)

References Connan et al. 1994; Menendez-Arias et al.

• Promotes assembly of HLA-A2 molecules in T2 cell lysates.

**HXB2 Location** RT (309-317)

Author Location RT (510-518)

**Epitope ILKEPVHGV** 

Immunogen in vitro stimulation or selection

Species (MHC) human (A2)

References Parker et al. 1992

• Studied in the context of HLA-A2 peptide binding.

**HXB2 Location** RT (309–317)

Author Location Pol (476–484)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Dyer et al. 1999

- CTL specific responses were measured over a 1.3 to 1.5 year period in members of the Sydney Blood Bank Cohort (SBBC) who had been infected with a natural attenuated strain of HIV-1 which was Nef-defective.
- · Some of these patients had prolonged high levels of CTL effector and memory cells despite low viral load.

**HXB2 Location** RT (309-317)

**Author Location** RT (476–484)

Epitope ILKEPVHGV

Immunogen in vitro stimulation or selection

Species (MHC) human (A2)

Keywords dendritic cells

References Zarling et al. 1999

· This study compares the ability of macrophages and dendritic cells to stimulate primary responses in CD8+ lymphocytes isolated from HLA-appropriate HIV-uninfected donors using peptide-pulsed APC – the dendritic cells performed better as APC for the stimulation of primary responses.

- Strong CTL responses were elicited by the epitopes DRFYK-TLRA and GEIYKRWII when presented by either immature or mature dendritic cells – macrophages were not able to prime a CTL response against DRFYKTLRA.
- A weak response to KLTPLCVSL was stimulated using macrophages as the APC.
- No detectable response was observed for the following previously-defined HIV epitopes: KIRLRPGGK, ILKEPVHGV, IRLRPGGK, GPKVKQWPL.

**HXB2 Location** RT (309–317) **Author Location** RT (480–)

Epitope ILKEPVHGV

Immunogen computer prediction

Species (MHC) (A2)

**Keywords** subtype comparisons **References** Schafer *et al.* 1998

- This study uses EpiMatrix for T cell epitope prediction to identify possible HLA-B27 and A-2 CTL epitopes in HIV.
- Based on EpiMatrix predictions, 28 peptides were synthesized and tested using T2 binding assays for potential HLA A2 or B27 binding, and 12 of these were shown to bind to the predicted HLA molecule.
- Two of these 12 peptides had been previously identified as CTL epitopes: HLA-B27 KRWILGLNK and HLA-A2 ILKEPVHGV
- This sequence is not conserved between clades, but is found only in a small number of B clade isolates.

**HXB2 Location** RT (309-317)

Author Location RT

**Epitope** ILKEPVHGV

**Epitope name** RT IV9

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** binding affinity, subtype comparisons, supertype, computational epitope prediction

References Altfeld et al. 2001c

- HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.
- Three additional previously described HLA-A2 epitopes were added to the set of 20, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- This peptide binds to four HLA-A2 supertype alleles: A\*0201, A\*0202, A\*0206 (highest affinity) and A\*6802.
- RT IV9 was recognized in 7/22 patients with chronic HIV-1 infection.
- 1/13 patients with acute HIV-1 infection recognized RT IV9.

**HXB2 Location** RT (309–317)

Author Location Pol (subtype A)

Epitope ILKDPVHGV

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** HIV exposed persistently seronegative (HEPS), escape

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- ILKDPVHGV or ILKEPVHGV was recognized in 1 of the 6 women (ML1760), and the response was present in the last available sample prior to seroconversion, 12 months.
- 20/20 sequences of the infecting strain had no substitutions in this epitope, all were ILKDPVHGV, so there was no evidence for escape.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized by 4/22 HEPS control sex workers: ML887, ML1192, ML1250, and ML1749.

**HXB2 Location** RT (309–317)

Author Location RT (476-484)

Epitope ILKEPVHGV

Epitope name ILK

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** HAART, ART, supervised treatment interruptions (STI), immunodominance, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- One of the 2/8 HLA-A2+ study subjects recognized this CTL epitope.
- Patient SC9 (HLA A1/2, B8/13, Cw0/0701, DR2/11, DQ6/7) had a CTL response against epitopes FLKEKGGL, ILKEPVHGV, SQRRQDILDLWIYHTQGYFPDWQNY, and GEIYKRWII peptides and all responses declined during therapy initiated at day 390 but were restored when therapy become intermittent.

**HXB2 Location** RT (309–317)

Author Location Pol

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Kostense et al. 2001

 HLA tetramers to six epitopes were used to study HLA-A2, B8 and B57 CTL in 54 patients – HIV-specific tetramer positive cells were inversely correlated with viral load in patients with high CD4, but in patients with CD4 T-cells below 400 high tetramer frequencies were found despite high viral load.

- sions, but many of these cells aren't functional.
- In 15 of the patients, the proportion of IFN gamma producing tetramer cells correlated with AIDS-free survival.

**HXB2 Location** RT (309–317)

**Author Location** Pol

**Epitope ILKEPVHGV** Immunogen HIV-1 infection Species (MHC) human (A2)

Keywords HAART, ART, immunodominance

References Seth et al. 2001

- CTL responses were studied by tetramer staining in 41 patients with combination therapy – activated CD8+ T-cells decline as the viral load drops in response to therapy, but the overall level of antigen-specific cells capable of differentiating into effectors stays constant and new epitopes may be recognized.
- 6/10 A\*0201+ individuals had HIV-specific tetramer staining cells, and 5 of these declined upon successful therapy.
- 3/10 A\*0201+ individuals with chronic HIV-1 infection recognized this epitope.
- Prior to therapy, the mean percentage of CD8+ cells that recognized theimmunodominant epitope SLYNTVATL was six-fold greater than the percentage recognizing the epitope ILKEPVHGV.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484 SF2)

**Epitope** ILKEPVHGV Immunogen HIV-1 infection

Species (MHC) human (A2)

Kevwords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A2+ individuals that had a CTL response to this epitope broken down by group: 1/10 group 1, 0/6 group 2, and 3/4 group 3.

**HXB2 Location** RT (309–317)

Author Location Pol (476–484)

Epitope ILKDPVHGV

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A2)

Keywords subtype comparisons, HIV exposed persistently seronegative (HEPS), immunodomi-

nance

References Kaul et al. 2001a

• Variants ILK(D/E)PVHGV are A/B clade specific.

- Most patients have high levels of HIV-specific T-cell expan- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
  - Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
  - 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
  - Among HLA-A2 women, 7/10 HEPS and 14/26 HIV-1 infected women recognized this epitope, and ILK(D/E)PVHGV tended to be more reactive in HEPS women, SL(F/Y)NTVATL in infected women.
  - The dominant response to this HLA allele was to this epitope in all 7/10 HEPS cases but in only 5 of the 14/26 HIV-1 infected
  - Four epitopes were considered to be "resistant epitopes", as they were preferentially reactive in HEPS women and so may confer resistance, and these were found in three different proteins: A2 ILK(D/E)PVHGV in RT, A\*6802 DTVLEDINL in Protease, B14 DLN(M/T)LN(I/V)V in p24 and B18 FRDYV-DRF(Y/F)K also in p24.
  - Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.
  - Subject ML 1250 had an A2 response to ILK(D/E)PVHGV prior to seroconversion, which switched to SL(F/Y)NTVATL post-seroconversion.
  - Subject ML 1760 had an A2 response to ILK(D/E)PVHGV prior to seroconversion, and gained responses to epitopes A2 SL(F/Y)NTVATL and B27 KRWII(L/M)GLNK postseroconversion.

**HXB2 Location** RT (309-317)

Author Location Pol (93TH253 subtype CRF01)

**Epitope** ILRIPVHGV Epitope name P464-472

Subtype CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers - weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not
- This epitope was reactive in HIV+ control study subject 144 who carried HLA-A2.

**HXB2 Location** RT (309-317)

Author Location Pol (93TH253 subtype CRF01)

Epitope ILRIPVHGV
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A2)
Konwords subtype comparis

**Keywords** subtype comparisons **References** Bond *et al.* 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- 1/4 tested FSWs recognized the E clade version of this epitope, which differs from the previously defined B clade version by two amino acids: ILKEPVHGV.
- This epitope was not conserved in many subtypes, and exact matches were very rare.

HXB2 Location RT (309–317)
Author Location RT (309–317)
Epitope ILKEPVHGV
Immunogen HIV-1 infection
Species (MHC) human (A2)

**Keywords** rate of progression, acute/early infection **References** Day *et al.* 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.

**HXB2 Location** RT (309–317)

**Author Location** 

Epitope ILKEPVHGV Epitope name Pol-IV9 Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2) References Sabbaj *et al.* 2003

• Among HIV+ individuals who carried HLA A02, 9/29 (31%) recognized this epitope.

HXB2 Location RT (309–317) Author Location Pol (476–484 LAI)

Epitope ILKEPVHGV

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2)

Keywords HAART, ART, epitope processing

References Kelleher et al. 2001a

Ritonavir (RTV) inhibits chymotryptic activity in the 20S proteasome *in vitro*, as does Saquinavir (SQV) to a lesser extent; Indinavir (IDV) does not. Thus there is concern protease inhibitors may adversely effect CTL epitope processing, but this paper indicates that processing is not inhibited at therapeutically relevant concentrations of RTV when the proteasome is functioning in in an intracellular context.

- RTV did not alter the presentation two RT A2 epitopes processed by distinct pathways: ILKEPVHGV, generated by the constitutive proteasome containing the MB1 beta subunit, and VIYQYMDDL which is dependent on IFNgamma induction of LMP7 which replaces MB1 in the immunoproteasome, and is destroyed by MB1 in the constitutive proteasome.
- RTV did not inhibit the processing and assembly of HLA-B35 or -A2, which are assembled with a rapid and moderate time course, respectively, or of HLA-A3, -B27 and -B39.

**HXB2 Location** RT (309–317)

**Author Location** Pol

Epitope ILKDPVHGV Immunogen HIV-1 infection Species (MHC) human (A2)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

**HXB2 Location** RT (309–317)

Author Location RT (476–484 NL43)

Epitope ILKEPVHGV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords class I down-regulation by Nef

References Yang et al. 2002

• Nef down-modulates class I protein expression, and this study demonstrates directly that Nef-deleted HIV-1 NL-43 can be more effectively killed *in vitro* than NL-43 with an intact Nef. The effect was shown to be specific for class I presentation of epitopes, and unlike Nef, deleting Vpr did not alter CTL susceptibility of NL-43 infected cells. The CTL clone 68A62, specific for the class I A2 presented ILKEPVHGV epitope, was one of four used in this study.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484 BRU)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Donor MHC A2

Keywords epitope processing

References Cohen et al. 2002

 The antigen presentation of two A2-restricted epitopes was compared, SLYNTVATL (p17) and ILKEPVHGV (RT). HIV-1 infected cells were more sensitive to lysis by SLYNTVATLspecific CTL than by ILKEPVHGV-specific CTL, because of a higher density of SLYNTVATL-A2 resulting from differences in processing.

- Incubation with a T1-cell proteolytic extract showed that by four hours, 25% of a p17 peptide had a C-term Leu-85 and were SLYNTVATL-precursors, while ILKEPVHGV-precursors were far less frequent (6.8%) even with four times more proteolytic extract after 30 hours.
- p17 was preferentially cleaved between Leu85 and Tyr86, while appropriate Val484 and Tyr485 cleavage was minor for RT.
- In a competition experiment, RSLYNTVATL bound TAP 3.7fold more efficiently than RT peptides.
- No difference in CTL avidity was detected in six patients with HLA-A2-restricted responses to these epitopes.
- No significant difference in HLA-A2 binding of to p17 or RT epitopes was observed.

HXB2 Location RT (309–317) Author Location Pol (476–484) Epitope ILKEPVHGV

Epitope name p9

Immunogen vaccine

*Vector/Type:* peptide *Adjuvant:* Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (A2)

References De Lucca et al. 2002

- BALB/c mice immunized with the p9 peptide, ILKEPVHGV, elicited specific lymphocyte proliferation activity.
- Exposure of lymphocytes from HIV-negative, HLA-A2 positive people to p9-RNA stimulated lymphocyte proliferation activity to p9. Anti-p9 CTL activity in human lymphocytes incubated with RNA extracted from lymphoid organs of p9-vaccinated mice could be more intensely stimulated.
- This murine RNA also mediated RNA-dependent protein kinase (PKR) and NFkappaB activation in the human lymphocytes, which may be driving the enhanced CTL stimulation in the human cells.

**HXB2 Location** RT (309–317)

**Author Location RT** 

Epitope ILKEPVHGV

Epitope name ILK

Immunogen HIV-1 infection Species (MHC) human (A2)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius *et al.* [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there was no correlation between CTL responses with viral rebound rates, plateau viral loads, or clearance rates.

**HXB2 Location** RT (309–317)

Author Location p51 (476-484)

Epitope ILKEPVHGV Immunogen vaccine

Strain: B clade IIIB HIV component: Gag,

Pol Adjuvant: IL-12

Species (MHC) mouse (A2) Donor MHC H2/Kb References Kmieciak et al. 2001

- Transgenic mice expressing a HLA-A2/Kb chimeric protein were vaccinated with either a p17-p24-p51 fusion protein (vG/P-92) or the Gag-Pol precursor protein (vVK1).
- Compared to vVK1, vG/P-92 induced a significant increase in Gag and Pol induced IFNgamma production and CTL responses, and to the epitopes SLYNTVATL and ILKEPVHGV, as determined by Elispot and 51Cr-release assays.

**HXB2 Location** RT (309–317)

Author Location RT (309–317 NL-43)

**Epitope** ILKEPVHGV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** class I down-regulation by Nef, escape **References** Ali *et al.* 2003

- NL43 was passaged in the presence of Nef KEKGGLEGL-specific CTL clones STD11 and KM3, giving rise to rapid selection of escape mutations, including E93G, E93K, K94N+G99R, G95R+G99R, E98K, E98D, G99R, G99E, L100P, and L100I; insertions, deletions, frameshifts and an early stop codon. 34/36 (94%) of sequences carried mutations in the epitope by seven days, 36/36 (100%) by 14 days.
- Several mutations selected by KEKGGLEGL-specific CTL were shown to impair the down regulation of class I MHC by Nef, in particular E93G, E93K, and a truncation mutation at position 51.
- Nef deletion mutants increased 100-fold NL-43 susceptibility to inhibition by CTL specific for epitopes in other proteins, the A2 epitopes ILKEPVHGV in RT and SLYNTVATL in p17 Gag.

**HXB2 Location** RT (309–317)

**Author Location** Pol (476–)

Epitope ILKEPVHGV

Epitope name Pol476

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This epitope was one of the previously identified HLA-A2 epitopes studied.
- 9/17 HIV-infected HLA-A2+ people recognized this epitope.

**HXB2 Location** RT (309–317)

Author Location RT (309-317)

Epitope ILKEPVHGV

**Epitope name** RT2

Subtype B

Immunogen vaccine, in vitro stimulation or selection

Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) transgenic mouse (A2) References Domingo et al. 2003

- A virus-like protein scaffold, called E2DISP, derived from pyruvate dehydrogenase multienzyme from Bacillus stearothermophilus has been engineered to display 60 copies of one or more epitopes on a single molecule.
- The E2DISP scaffold displaying pep23 is able to stimulate a Th responses, and peptide RT2, which is a CTL epitope from HIV-1 RT, was able to elicit a CD8+ T cell response in vitro and in a vaccinated HLA-A2 transgenic mouse. Thus the E2DISP scaffold allows cell-entry and access to the class I and class II processing pathways.

**HXB2 Location** RT (309–317)

**Author Location** Pol (476–484)

**Epitope ILKEPVHGV** Immunogen HIV-1 infection Species (MHC) human (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Tetramer binding,

Flow cytometric T-cell cytokine assay

Keywords responses in children References Sandberg et al. 2003

- 65 vertically HIV-1 infected children, ages 1-16, the majority undergoing ART, were analyzed in regard to their plasma viremia and CD4+ and CD8+ T cell counts, and CD8+ T cell responses.
- Using vaccina expressed Gag, Pol, Env, Rev, Nef in target cells in an Elispot assay, 85% of the children recognized at least one HIV antigen. The strong CD8+ T cell responses were directed against Pol, followed by Gag and Nef. Children younger than 4 had significantly weaker responses (7/14 had no reponse) than older children (only 1/32 had no response, and responses were greater in magnitude).
- · SLYNTVATL and ILKEPVHGV tetramers were used to quantitate specific responses. 49 chlidren in an expanded cohort carried HLA-A2. 1/11 children under 3 years of age had detectable CD8+ T-cell responses to SLYNTVATL, 2/11 to ILKEPVHGV. Among children over 3, 11/38 recognized SLYNTVATL and 9/38 recognized ILKEPVHGV.
- Older children that maintained a CD4 count greater than 400 cells/ul tended to have stronger CTL reponses.

**HXB2 Location** RT (309–317)

**Author Location** RT (309–317)

**Epitope ILKEPVHGV** 

**Immunogen** HIV-1 infection

Species (MHC) (A2)

**Donor MHC** A2, A3, B27, B51; A2, A3, B27, B57; A2, A23, B57

Assay type Cytokine production, CD8 T-cell Elispot -IFNγ, Tetramer binding, Intracellular cyto-

kine staining **Keywords** assay standardization/improvement, memory

cells

References Sun et al. 2003

- Vector/Type: peptide HIV component: RT This study compares assay methods for testing CTL responses using samples from 20 HIV+ patients. The study compares ELISpot, tetramer-binding, and intracellular IFNy. Tetramerbidning analysis was performed with Gag (SLYNTVATL) or Pol (ILKEPVHGV) tetramers. Antigen presentation using recombinant vaccinia viruses (rVVs) encoding HIV-LAI Gag, Pol, Env, Nef, Tat and Vif proteins was compared to peptide panels. HIV antigen recognition in memory CTLs was measured by chronium release assay and compared to effector/memory CD8+ T cells in an IFN-γ ELISpot assay.
  - Results: IFNγ Elispot and flow cytometry gave similar frequencies of HIV specific CD8+ Tcells. Tetramer-binding analysis was most sensitive. Pools of peptides and the sum of frequencies of individual peptides were comparable. Elispot assays using peptides were more sensitive than assays using vaccinia expressed proteins. Cr release and Elispot against rVVs gave comparable memory cell responses 2/3s of the time.
  - 3/7 HLA-A2+ patients recognized this epitope.

**HXB2 Location** RT (309-317)

**Author Location** RT (309–317 NL43)

**Epitope ILKEPVHGV** 

Epitope name IV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type Chromium-release assay, CTL suppression of replication

Keywords escape

References Yang et al. 2003a

- · Virus was cultured in the presence of CTL lines specific for 4 different epitopes (SLYNTVATL, ILKEPVHGV, IEIKDTKEAL, and KEKGGLEGL) to study the emergence of escape mutations. Escape varied between clones for the same epitope, and between different epitopes. Gag and RT epitope escape, if it occurred at all, tended to be monoclonal and within the epitope, indicating strong fitness constraints, while the Nef epitope escape was rapid, polyconal, and sometimes the result of upstream frameshifts.
- There was one cloned cell line that recognized ILKEPVHGV, 68A62. After 2 weeks of passaging HIV-1 in the presence of 68A62, the mutated epitope ilkeLvhgv was found in 6/12 sequences.

**HXB2 Location** RT (309-317)

**Author Location** Pol

**Epitope** ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

Keywords HIV exposed persistently seronegative (HEPS)

References Koning et al. 2004

• A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.

 2/11 HLA A2+ infection-resistant men, compared to 1/9 preseroconversion men who went on to become infected, reacted to this epitope.

**HXB2 Location** RT (309–317)

**Author Location** RT Pol (464–472)

**Epitope ILKEPVHGV** 

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses.
   HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 9/19 patients recognized this epitope.

**HXB2 Location** RT (309–317)

**Author Location** RT (309–317)

Epitope ILKEPVHGV

Epitope name RT-IV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type Chromium-release assay

Keywords binding affinity, TCR usage, characterizing

CD8+ T cells

References Yang et al. 2003b

- Killing efficiency of CTLs and their ability to suppress viral replication is shown to depend more on epitope specificity than to antigenic avidity. Different clones recognizing the same epitope had similar killing efficiency despite their variation in avidity. Nef specific CTL clones tended to be most inhibitory, followed by Gag, then by RT specific clones, regardless of avidity.
- 2/14 CTL T-cell clones tested were specific for RT/IV9. Under conditions of excess peptide (100ug/ml), there was no difference in their lytic potential; all possessed similar effector capacity. Avidity was measured as the sensitizing dose of peptide required for 50% of maximal killing (SD50), which varied from 20 pg/ml to 100 ng/ml, over four orders of magnitude for all 14 epitopes. The SD50 values for the two RT/IV9 clones were very different, 50 and 20,000 pg/ml.

**HXB2 Location** RT (309–317)

Author Location (B consensus)

Epitope ILKEPVHGV

Epitope name IV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A02, A03, B08, B62, Cw7, Cw10

Country United States.

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** RT (309-317)

Author Location RT (309-317)

**Epitope** ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type Chromium-release assay

Keywords assay standardization/improvement

References Lubong et al. 2004

 Using IL7 or IL15 in culturing of HIV-1-specific CTL clones was inferior to using IL-2 alone; the addition of these cytokines to IL-2 did not show any advantage. Neither proliferation, survival, nor lytic capacity of HIV-1-specific CTLs was significantly enhanced by addition of IL7 or IL15.

**HXB2 Location** RT (309-317)

**Author Location** Pol

Epitope ILKEPVHGV

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2)

**Donor MHC** A\*02, A\*30, B\*4402, B\*15

Assay type Tetramer binding, T-cell Elispot

**Keywords** HIV exposed persistently seronegative (HEPS)

References Missale et al. 2004

- HIV-specific T-cell response was tested in patients exposed to blood from a patient with highly replicating HIV; these patients were nosocomially infected with HBV, but uninfected with HIV. HIV-specific T-cell responses were directed to structural and non-structural HIV proteins in 2 patients, suggesting that the virus replicated in these patients sufficiently to prime a cell-mediated immune response that protected them from HIV infection.
- This patient responded to 4/8 HIV epitopes tested in an IFNgamma EliSpot assay or tetramer assay. Responses were detected 8 and 28 weeks after exposure. A response to ILKEPVHGV was detected by both assays.

**HXB2 Location** RT (309–317)

**Author Location** Pol

Epitope ILKEPVHGV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United Kingdom.

Assay type Tetramer binding, T-cell Elispot, Intracellular

cytokine staining

**Keywords** rate of progression, acute/early infection, char-

acterizing CD8+ T cells, immune dysfunction

References Papagno et al. 2004

Acute HIV-1 infection induces massive activation of HIV-specific and non-HIV-specific CD8+ T-cells resulting in differentiation of these cells. High differentiation of CD8+ T-cells is correlated with disease progression. Differentiation is a natural process but it can be driven by elevated immune activation, such as in HIV infection.

**HXB2 Location** RT (309–317)

Author Location (309–317)

Epitope ILKEPVHGV

Epitope name RT2

Immunogen vaccine

Vector/Type: bacteriophage coat protein, dihydrolipoyl acetyltransferase E2 protein, of Bacillus stearothermophilus HIV compo-

nent: RT

Species (MHC) transgenic mouse (A2)

**Assay type** Chromium-release assay **Keywords** vaccine antigen design

**References** De Berardinis *et al.* 2003

An RT T-helper (KDSWTVNDIQKLVGK) that can be promiscuously presented by multiple HLA-DR molecules, and an RT CTL epitope (ILKEPVHGV) presented by HLA-A2, were displayed using two different antigen presentation systems, bacteriophage virions or E2 protein scaffolds. Both systems enabled display of the epitopes in a mouse model system to the immune system. CTL responses were detected in immunized mice, and were processed correctly for both class I and class II

**HXB2 Location** RT (309–317)

**Author Location** Pol

presentation.

Epitope ILKEPVHGV

Epitope name IV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Two escape mutations, at positions 3 ilRepvhgv, and 9 ilkepvhgA, were found not to correspond to the most polymorphic residues in the epitope.

**HXB2 Location** RT (309–317)

**Author Location** RT (309–317)

Epitope ILKEPVHGV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

- The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection.
- ILKEPVHGV was targeted in 54% of 74 A2+ chronically infected individuals, but only 1/14 acutely infected A2+ individuals

**HXB2 Location** RT (309-317)

**Author Location** 

**Epitope** ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Germany.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, escape, variant cross-

recognition or cross-neutralization, optimal

epitope

References Harrer et al. 2005

- An HLA-B13-restricted optimal epitope was defined in Nef, RI9. The frequency of CTLs specific for this epitope in B13positive patients exceeded the number of CTLs against other epitopes, indicating that this is a dominant epitope in B13positive subjects. Three B13-positive patients who had an immunodominant response to this epitope were good controllers of their infection, with low viral loads over long periods.
- 5 HLA A2+ B13+ patients were found to make an immunodominant response to the B13 epitope RI9. 0/5 recognized the A2 epitope ILKEPVHGV, and only 1/5 recognized the A2 epitope SLYNTAVTL, with a much lower frequency than the B13 response.

**HXB2 Location** RT (309–317)

Author Location Pol (476–484 BRU)

Epitope ILKEPVHGV

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 2/9 CRF02\_AG-infected Ivorians, and 1/9 B-infected French subjects.

**HXB2 Location** RT (309–317)

**Author Location** RT (77–85)

Epitope ILKEPVHGV

Epitope name IL9

Subtype B

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A2)

Country Canada.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance, genital and mucosal immunity, characterizing CD8+ T cells

References Makedonas et al. 2005

- CD8 T-cell responses were studied in individuals who remained seronegative in spite of being mucosally (group 1) or intravenously (group 2) exposed to HIV-1. A similar proportion of subjects from each group recognized at least 1 HIV peptide, and they recognized peptides with similar cumulative intensity. The proportion of responding individuals in both groups was significantly greater than in a low-risk, negative control group. One exposed uninfected subject recognized 7 epitopes.
- HLA-A\*0201 epitopes that are immunodominant in chronically infected individuals were rarely stimulatory in exposed uninfected individuals. SLYNTVATL was recognized by one HLA A2+ individual in each group (1/11 vs 1/5), while none of the exposed uninfected individuals tested responded to ILKEPVHGV. In contrast, chronically infected subjects recognized these epitopes at a frequency of 69% and 31%, respectively.

**HXB2 Location** RT (309–317)

**Author Location** RT (476–484)

**Epitope** ILKEPVHGV

Epitope name IV9

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Germany.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** HAART, ART, TCR usage, characterizing CD8+ T cells, optimal epitope

References Schmitt-Haendle et al. 2005

- CTL responses to 3 HLA-A2-restricted epitopes were investigated in 51 HIV-1 infected HLA-A2+ individuals. The most prevalent response was seen for IV9, followed by SL9. The VL9 epitope was not recognized. There was a significant correlation of CTL activity to the CD8 counts in peripheral blood, but no correlation to CD4 counts, viral load, or antiviral therapy
- 37.3% of the individuals recognized ILKEPVHGV.
- All analyzed mutations for RT-IV9 epitope could decrease or abrogate CTL recognition dependent on the CTL clones tested, but all were fully immunogenic for other CTL clones. The ilkDpvhgv, ilkepvhEv, Rlkepvhgv and ilRepvhgv variants were tested.

**HXB2 Location** RT (309–317)

Author Location Pol (476–484)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind three of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

**HXB2 Location** RT (309–317)

**Author Location** Pol (464–472)

Epitope ILKEPVHGV

Immunogen HIV-1 infection

Species (MHC) human (A2, A\*0201)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** RT (309–317)

**Author Location** Pol (subtype B)

**Epitope** ILKEPVHGV

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2, A\*0202)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among B and D clade viruses.
- Clade A version of the epitope, ILKDPVHGV, was preferentially recognized by CTL.

**HXB2 Location** RT (309–317)

Author Location RT (309-317)

Epitope ILKEPVHGV

Epitope name RT2

Immunogen vaccine, in vitro stimulation or selection

Vector/Type: HIV-1 peptide in filamentous bacteriophage major coat protein HIV com-

ponent: RT

Species (MHC) human, mouse (A2, A2 transgenic)

Keywords epitope processing

References De Berardinis et al. 2000

 Phage display of the CTL epitope, ILKEPVHGV coupled with T helper epitope KDSWTVNDIQKLVGK, elicited specific CTL responses in vitro in PBMC from HIV negative individuals in and in vivo in immunization of HLA-A2 transgenic mice. • Bacteriophage presentation of peptides is generally used for stimulation of antibodies, and this novel discovery of CTL epitope processing and presentation suggests new possibilities for these vectors.

**HXB2 Location** RT (309–317) **Author Location** RT (476–484 LAI) Epitope ILKEPVHGV Subtype B Immunogen HIV-1 infection Species (MHC) human Donor MHC A\*0201

Keywords HAART, ART, responses in children

References Luzuriaga et al. 2000

- Longitudinal study of 8 infants with prolonged viral suppression due to combination antiretroviral therapy showed no HIV-1 specific CTL responses in peripheral blood cells. 6/8 were studied using a Chromium release assay and no response was detected using Gag expressed in vaccinia in the target cells. Three HLA-A\*0201 children were tested using SLYNTVATL or ILKEPVHGV HLA A\*0201 tetramers and again no HIVspecific response was detected, either using PBMC specimens, or PBMC which had been stimulated in vitro for a week.
- In contrast, one of the children with suppressed HIV viral replication who was co-infected with HIV and EBV, while HIV-tetramer negative, had EBV-tetramer staining cells at a frequency of 0.14% in the PBMC.

**HXB2 Location** RT (309-318)

**Author Location** Pol

**Epitope** ILKEPVHGVY

Epitope name 1249

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A2)

Donor MHC A02, A30, B39; A02, A03, B44, Cw05, • Review of HIV CTL epitopes.

Cw07; A02, A30, B35, B49, Cw04, Cw07

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope prediction, cross-presentation by different HLA

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for ILKEPVHGVY: 96% Promiscuous epitope binding to A02 and Bw62.

**HXB2 Location** RT (309–318) **Author Location** RT (476–485 LAI) **Epitope ILKEPVHGVY** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1501)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1501 epitope.

**HXB2 Location** RT (309-318)

Author Location RT (309-317)

Epitope ILKEPVHGVY

Immunogen HIV-1 infection

Species (MHC) human (B15)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

HXB2 Location RT (309-318)

**Author Location** RT (476–485 LAI)

**Epitope** ILKEPVHGVY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B62)

Kevwords review

References McMichael & Walker 1994: Menendez-Arias

et al. 1998

**HXB2 Location** RT (309-318)

Author Location RT (309-318)

Epitope IKLEPVHGVY

Immunogen HIV-1 infection

Species (MHC) human (B62)

Keywords immunodominance

References Day et al. 2001

• No immunodominant responses were detected to four B62restricted epitopes tested.

HXB2 Location RT (309-318)

Author Location Pol

**Epitope ILKEPVHGVY** 

Subtype A, B, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade

HIV component: p17 Gag, p24 Gag

Species (MHC) human (B62)

Keywords subtype comparisons, epitope processing,

vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location RT (317–325)
Author Location Pol (484–492)
Epitope VYYDPSKDL
Subtype B, CRF02\_AG
Immunogen HIV-1 infection
Species (MHC) human (A24)
Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, immunodominance, characterizing CD8+ T cells

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 1/9 CRF02\_AG-infected Ivorians, and 0/9 B-infected French subjects.

**HXB2 Location** RT (328–352) **Author Location** RT (495–515 LAI)

Epitope EIQKQGQGQWTYQIYQEPFKNLKTG

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A11)

**References** Menendez-Arias *et al.* 1998; Walker *et al.* 1989

• One of five epitopes defined for RT-specific CTL clones in this study.

**HXB2 Location** RT (333–341)

**Author Location** 

Epitope GQGQWTYQI

**Epitope name** GI9

Immunogen

Species (MHC) human (B13)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B13 epitope.

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HXB2 Location RT (340–350)
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**Author Location** Pol (487–497 93TH253 subtype CRF01)

Epitope QIYQEPFKNLK Epitope name P495-505 Subtype CRF01\_AE

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was weakly reactive in the HEPS study subject 128 who was HLA A11/A33.
- This epitope was reactive in HIV+ study subjects 053 and 184 who carried HLA-A11.

**HXB2 Location** RT (340–350)

Author Location Pol (487–497 93TH253 subtype CRF01)

Epitope QIYQEPFKNLK
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A11)

**Keywords** subtype comparisons **References** Bond *et al.* 2001

- HLA-A11 CRF01 (called subtype E in Bond *et al.*) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This epitope was not predicted by the EpiMatrix method to be likely to bind to A11, though it served as an epitope in the FSWs, and it was one of the six A11 epitopes that had been previously defined.
- 5/8 tested FSWs recognized this epitope.
- This epitope was highly conserved in other subtypes, although exact matches were not very common.

**HXB2 Location** RT (340–350)

**Author Location** RT (507–516)

Epitope QIYQEPFKNLK

Immunogen HIV-1 infection

Species (MHC) human

References Menendez-Arias et al. 1998; Price et al. 1995

• Study of cytokines released by HIV-1 specific activated CTL.

HXB2 Location RT (340–352) Author Location RT (507–519 LAI) Epitope QIYQEPFKNLKTG Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

Kevwords review

**References** Johnson & Walker 1994; Menendez-Arias *et al.* 1998

• This epitope was listed in a review.

**HXB2 Location** RT (340–352)

Author Location Pol (495-507)

**Epitope** QIYQEPFKNLKTG **Immunogen** HIV-1 infection

Species (MHC) human (A11)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** RT (341–349)

**Author Location** (C consensus)

Epitope IYQEPFKNL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** RT (341–349)

Author Location (C consensus)

Epitope IYQEPFKNL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- IYQEPFKNL is an optimal epitope for both A\*2301 and A\*2402.

**HXB2 Location** RT (341–349)

**Author Location** (C consensus)

**Epitope IYQEPFKNL** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- IYQEPFKNL is an optimal epitope for both A\*2301 and A\*2402.

**HXB2 Location** RT (341–350)

Author Location RT (508-516)

Epitope IYQEPFKNLK

Immunogen HIV-1 infection

Species (MHC) human (A\*1101)

References Culmann 1998

 C. Brander notes that this is an A\*1101 epitope in the 1999 database.

**HXB2 Location** RT (341–350)

**Author Location** RT (508–517 LAI)

Epitope IYQEPFKNLK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*1101)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*1101 epitope.

**HXB2 Location** RT (341–350)

Author Location (C consensus)

Epitope IYQEPFKNLK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*1101)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- IYQEPFKNLK is an optimal epitope.

**HXB2 Location** RT (341–350)

**Author Location** RT (508–517 SF2)

Epitope IYQEPFKNLK

**Immunogen** HIV-1 infection

Species (MHC) human (A11)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A11+ individuals that had a CTL response to this epitope broken down by group: 0/3 group 1, 0/0 group 2, and 1/2 group 3.

HXB2 Location RT (341–350) Author Location Pol (508–516) Epitope IYQEPFKNLK

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

HXB2 Location RT (341–350)
Author Location Pol (497–506)
Epitope IYQEPFKNLK
Epitope name IK10
Subtype B
Immunogen HIV-1 infection

Species (MHC) human (A11)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

**Keywords** optimal epitope **References** Allen *et al.* 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- This epitope did not vary.

HXB2 Location RT (341–350) Author Location Pol (497–506) Epitope IYQEPFKNLK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A11)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** RT (349–366)

Author Location (C consensus)

Epitope LKTGKYAKMRTAHTNDVK

Subtype C

Immunogen HIV-1 infection Species (MHC) human (Cw\*0602) Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (356–365)

**Author Location** 

Epitope RMRGAHTNDV

Epitope name Pol-RV10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*3002)

**Donor MHC** A\*2904 A\*3002 B\*1503 B\*5802 Cw\*0202 Cw\*0602

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- Subject 01RCH50 also recognized the epitope WRFDSRLAF, Nef(183-191), B\*1503.
- Among HIV+ individuals who carried HLA A30, 5/16 (31%) recognized this epitope.

HXB2 Location RT (356–365) Author Location RT (356–365)

Epitope RMRGAHTNDV

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** RT (356–366)

Author Location RT (356–366)
Epitope RMRGAHTNDVK
Immunogen HIV-1 infection
Species (MHC) human (A\*0301)
Keywords optimal epitope
References Frahm et al. 2007

HXB2 Location RT (356–366)
Author Location RT (15–26)
Epitope RMRGAHTNDVK
Epitope name A3-RK11
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (A3)
Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 5/7 individuals began to have detectable responses to this epitope after STI.

Author Location RT (356–366)

Author Location RT (356–366)

Epitope RTRGAHTNDVK

Epitope name A3-RK11 Pol

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

- An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.
- The second infecting strain had the variant rtrgahtndvR. The CTL response to both variants declined over time, and the response to the second variant was lower than to the first throughout.

HXB2 Location RT (356–366)
Author Location (B consensus)
Epitope RMRGAHTNDVK
Epitope name RK11
Subtype B
Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A02, A03, B08, B62, Cw7, Cw10

Country United States.

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

HXB2 Location RT (356–366) Author Location Pol (512–522) Epitope RMRGAHTNDVK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** RT (364–372)

**Author Location** RT (518–526 U455)

Epitope DVKQLTEVV

Immunogen

**Species (MHC)** human (A28, A\*6802)

**Keywords** subtype comparisons

References Dong 1998; Menendez-Arias et al. 1998

- Predicted on binding motif, no truncations analyzed.
- Reacts with clade A consensus (U455), and with the peptide DVKQLAEAV, from the D clade.

**HXB2 Location** RT (364–372)

**Author Location** RT (470–478 subtype A)

**Epitope** DVKQLTEVV

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (B70)

**Keywords** subtype comparisons

References Dorrell et al. 1999

• CTL responses in three individuals with non-clade B infections were studied, 2 with subtype A infections, 1 with subtype C – their infections all originated in East Africa.

- This CTL response was defined in a patient with an A subtype infection.
- Bulk cultures from this patient gave a CTL response that could recognize the subtype D form of this epitope, with two substitutions (DVKQLAEAV), though a CTL line from these cultures didn't recognize the B clade variant (DVKQLTEAV)

HXB2 Location RT (366–385) Author Location Pol (521–540)

Epitope KQLTEAVOKIAMESIVIWGK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** RT (373–390)

Author Location RT (373–390 HXB2)

Epitope QKIATESIVIWGKTPKFK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 21% of the study subjects, and it was one of the 25 most frequently recognized peptides.

HXB2 Location RT (374–383) Author Location RT (LAI) Epitope KITTESIVIW Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords rate of progression

**References** Menendez-Arias *et al.* 1998; van der Burg *et al.* 1997

- Patients studied were from the Amsterdam cohort.
- CTL epitopes of 3 rapid progressors were compared to 4 long-term survivors (LTS); no differences could be found in the degree of conservation between them.
- Epitope recognized by LTS and by a progressor.

**HXB2 Location** RT (374–383)

Author Location RT (LAI)

Epitope KITTESIVIW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

References van der Burg et al. 1997

 Recognized by CTL from a progressor and a long-term survivor, PIVLPEKDSW was also recognized.

**HXB2 Location** RT (374–383)

**Author Location** RT Pol (529–538)

Epitope KITTESIVIW

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 7 patients recognized this epitope.

**HXB2 Location** RT (375–383)

**Author Location** RT (375–383 LAI)

Epitope ITTESIVIW

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5701, B\*5801)

Keywords rate of progression

References Klein et al. 1998

- Another patient recognized the ten-mer version of this epitope, KITTESIVIW van der Burg et al. [1997]
- B57 has been associated with long-term non-progression in the Amsterdam cohort.
- The most pronounced CTL responses in HLA B\*5701 LTS were to RT and Gag.
- The patient that recognized ITTESIVIW also recognized IVLPEKDSW.

**HXB2 Location** RT (375–383) **Author Location** (C consensus)

**Epitope** IAMESIVIW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5703)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- IAMESIVIW is an optimal epitope for both B\*5801 and B\*5703

**HXB2 Location** RT (375–383)

**Author Location** RT (375–383)

**Epitope IAMESIVIW** 

Immunogen

Species (MHC) human (B\*5801)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** RT (375–383)

**Author Location** (C consensus)

**Epitope** IAMESIVIW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the S5 residue of IAMESIVIW are associated with the presence of the HLA presenting molecule in the host.
- IAMESIVIW is an optimal epitope for both B\*5801 and B\*5703

**HXB2 Location** RT (375–383)

Author Location (C consensus)

Epitope IAMESIVIW

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*5801, B\*57)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords cross-presentation by different HLA, charac-

terizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** RT (375–383)

Author Location RT (375–383 SF2)

**Epitope ITTESIVIW** 

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B57+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 0/0 group 2, and 1/2 group 3.

**HXB2 Location** RT (392–401)

**Author Location** RT (559–568 LAI)

**Epitope** PIQKETWETW

Subtype B

Immunogen

Species (MHC) human (A\*3201)

**References** Harrer *et al.* 1996b; Menendez-Arias *et al.* 1998

- Reviewed in Menendez-Arias et al. [1998], suggest the epitope is HLA B53/Cw2.
- C. Brander notes that this is an A\*3201 epitope in the 1999 database.

**HXB2 Location** RT (392–401)

Author Location RT (559–568 LAI)

Epitope PIQKETWETW

Subtype B

Immunogen

Species (MHC) human (A\*3201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*3201 epitope.

**HXB2 Location** RT (392-401)

**Author Location** 

Epitope PIQKETWETW

Epitope name Pol-PW10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*3201)

**Donor MHC** 01RCH59: A\*0201, A\*3201, B\*4002,

B\*5301, Cw\*0202, Cw\*0401

**Keywords** HAART, ART **References** Sabbaj *et al.* 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previou.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated.
- Subject 01RCH59 was Hispanic, was not on HAART, viral load 5100, CD4 count 349, and she also recognized QASQEVKNW, p24(176-184), B\*5301.
- Among HIV+ individuals who carried HLA A32, 1/2 (50%) recognized this epitope.

**HXB2 Location** RT (392-401)

**Author Location** RT (559–568 SF2)

Epitope PIQKETWETW

Immunogen HIV-1 infection

Species (MHC) human (A32)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A32+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 1/1 group 2, and 0/0 group 3.

**HXB2 Location** RT (392–401)

**Author Location RT** 

**Epitope** PIQKETWETW

**Epitope name** A32-PW10(RT)

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A32)

Donor MHC A32, B7, B14; A32, B44; A30, A32, B18,

B27

Keywords HAART, ART, supervised treatment interrup-

tions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals. Patient A displayed the greatest response to epitope B14-EL9(gp41), a strong response to B7-TL9(p24), and responses to B7-TM9(Nef) and A32-PW10(RT). Patient B displayed the greatest response to epitope B44-AW11(p24) and also responded to A32-PW10(RT) in both PB and LN samples, while a third response against epitope A32-RW10(gp120) was only detected in the LN sample. Patient D displayed the greatest response to B27-KK10 (p24), and also responded to A30-RY11(p17), A32-PW10(RT), A30-KY11(RT), A32-RW10(gp120), and B18-YY9(Nef).

**HXB2 Location** RT (397–406)

**Author Location** RT (LAI)

**Epitope** TWETWWTEYW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B44)

**References** Menendez-Arias *et al.* 1998; van der Burg *et al.* 1997

 Recognized by CTL from two progressors, EILKEPVGHGV and EELRQHLLRW were also recognized by one, and RETKL-GKAGY was also recognized by the other.

**HXB2 Location** RT (397–406)

Author Location RT Pol (552–561)

Epitope TWETWWTEYW

Immunogen HIV-1 infection

Species (MHC) human (B44)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

• Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase

lated with viral load.

• Less than 2 of 11 patients recognized this epitope.

**HXB2 Location** RT (407–416) **Author Location** (C consensus)

Epitope QATWIPEWEF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- QATWIPEWEF is an optimal epitope for both B\*5702 and B\*5703.

**HXB2 Location** RT (407–416)

**Author Location** (C consensus)

**Epitope QATWIPEWEF** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5703)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- OATWIPEWEF is an optimal epitope for both B\*5702 and B\*5703.

**HXB2 Location** RT (407–416)

**Author Location** (C consensus)

Epitope QATWIPEWEF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

• HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.

significantly until the end of the follow up, but were not corre
• This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** RT (413–429)

Author Location (C consensus)

Epitope EWEFVNRPPLVKLWYQL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*8101)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression

References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (416-423)

Author Location Pol (571–)

**Epitope** FVNTPPLV

Epitope name Pol571

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: RT Adjuvant: Incomplete Freund's Adjuvant

(IFA)

**Species (MHC)** transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced a CTL responses 1/6 transgenic mice. Responses to the peptide were not detected in 17 HIV+ HLA-A2 subjects.

**HXB2 Location** RT (416–424)

**Author Location** Pol (563–571 93TH253 subtype CRF01)

**Epitope** FVNTPPLVK

Epitope name P571-579

Subtype CRF01\_AE

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A11)

Keywords HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

• This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.

- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was weakly reactive in the HEPS study subject 128 who was HLA A11/A33.

**HXB2 Location** RT (416–424)

Author Location Pol (563–571 93TH253 subtype CRF01)

Epitope FVNTPPLVK
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A11)
Keywords subtype comparisons

References Bond et al. 2001

- HLA-A11 CRF01 (called subtype E in Bond *et al.*) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This is one of the new A11 epitopes identified through the streamlined EpiMatrix method, and 1/8 tested FSWs recognized it.
- This epitope was conserved many subtypes (but not subtype H), but exact matches were not very common.

**HXB2 Location** RT (421–429)

**Author Location** Pol

Epitope PLVKLWYQL

Epitope name P9L Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV component: gp140, gp140ΔV3

Species (MHC) transgenic mouse (A\*0201)

**Assay type** Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted
CTL responses against multiple conserved HIV-1 epitopes in
HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw
virus expressing gp140 with deleted V1, V2, and V3 loops
successfully induces neutralizing antibodies against HIV-1. A
live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination
against HIV and measles.

**HXB2 Location** RT (421–429)

Author Location RT (421–429)

Epitope PLVKLWYQL
Immunogen HIV-1 infection
Species (MHC) human (A2)

References Haas et al. 1998

- Of 98 patients in cross-sectional analysis, 78% had CTL against pol – RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)
- New clusters of epitopes were defined utilizing different HLA molecules.

**HXB2 Location** RT (432–440)

Author Location RT (587-597 SF2)

Epitope EPIVGAETF Immunogen HIV-1 infection Species (MHC) human (B\*3501)

Keywords review

**References** Menendez-Arias *et al.* 1998; Tomiyama *et al.* 1997

- A CTL clone responsive to this epitope was obtained.
- 5/7 B35-positive individuals had a CTL response to this epitope.
- An E to D substitution at position 1, and V to I at position 4, reduces activity but not binding to B\*3501.
- Menendez-Arias *et al.* [1998] note in their review that this epitope is near the protease cleavage site and conservation of this region is important for proper viral maturation.

**HXB2 Location** RT (432–440)

**Author Location** Pol (587–595)

Epitope EPIVGAETF

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

References Tomiyama et al. 2000a

- CD8+ T-cells that bound one of six HIV-specific B\*3501epitope tetramers did not express CD28 or CD45A.
- A significant increase in CD28-CD45RA- cells and a decrease of CD28+CD45RA+ cells was observed in chronically HIV-1infected individuals relative to healthy individuals.
- CD28-CD45RA- cells are likely to be effector cells and have high levels of perforin in their cytoplasm.
- The mean percentage of total CD28- CD8+ cells in chronically infected HIV-1-infected patients was 76.6% in comparison to HIV-1-uninfected individuals (40.6%)

HXB2 Location RT (432-440)

**Author Location** 

Epitope EPIVGAETF

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords acute/early infection

References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.

- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

Author Location RT (432–440)
Author Location Pol (587–595)
Epitope EPIVGAETF
Immunogen HIV-1 infection
Species (MHC) human (B35)
References Dyer et al. 1999

- CTL specific responses were measured over a 1.3 to 1.5 year period in members of the Sydney Blood Bank Cohort (SBBC) who had been infected with a natural attenuated strain of HIV-1 which was Nef-defective.
- Some of these patients had prolonged high levels of CTL effector and memory cells despite low viral load.

**HXB2 Location** RT (432–440)

Author Location Pol

Epitope EPIVGAETF

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** HIV exposed persistently seronegative (HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- No one, 0/3 HLA B35+ infection-resistant men, and 0/5 preseroconversion men who went on to become infected, reacted to this epitope.

**HXB2 Location** RT (432–440)

Author Location RT (587-596 SF2)

Epitope EPIVGAETF

Immunogen HIV-1 infection

Species (MHC) human (B35, B51)

References Shiga et al. 1996

 Binds HLA-B\*3501, and is also presented by B51 – but CTL could not kill RT-vaccinia virus infected cells that expressed B51. HXB2 Location RT (432-440)

Author Location Pol (587-595)

**Epitope** EPIVGAETF

Immunogen HIV-1 infection

Species (MHC) human (B35, B51)

References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** RT (432–440)

Author Location RT (432-440)

Epitope EPIVGAETF

Immunogen HIV-1 infection

Species (MHC) human (B51)

**Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

**HXB2 Location** RT (432–441)

Author Location Pol (587–596)

**Epitope** EPIVGAETFY

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

References Tomiyama et al. 2000a

- CD8+ T-cells that bound one of six HIV-specific B\*3501-epitope tetramers did not express CD28 or CD45A.
- A significant increase in CD28-CD45RA- cells and a decrease of CD28+CD45RA+ cells was observed in chronically HIV-1infected individuals relative to healthy individuals.
- CD28-CD45RA- cells are likely to be effector cells and have high levels of perforin in their cytoplasm.
- The mean percentage of total CD28- CD8+ cells in chronically infected HIV-1-infected patients was 76.6% in comparison to HIV-1-uninfected individuals (40.6%)

**HXB2 Location** RT (432–441)

**Author Location** RT (587–597 SF2)

Epitope EPIVGAETFY

Immunogen HIV-1 infection

Species (MHC) mouse (B35)

Keywords review

References Menendez-Arias et al. 1998; Shiga et al. 1996

• Binds HLA-B\*3501, but not presented by B51, in contrast to the peptide EPIVGAETF.

- Menendez-Arias et al. [1998] note in their review that this epitope is located near the protease cleavage site and conservation of this region is important for viral maturation.
- This epitope spans the Pol p66 RT p15 (RNAse) domain.

HXB2 Location RT (432–441)
Author Location RT (587–597 SF2)
Epitope EPIVGAETFY
Immunogen HIV-1 infection

Species (MHC) human (B35)

**Keywords** rate of progression **References** Kawana *et al.* 1999

• HLA B35 is associated with rapid disease progression.

- The sequences of 9 previously described HIV-1 B35 CTL epitopes were obtained in 10 HLA B35+ and 19 HLA B35- individuals.
- 3/9 CTL epitopes had substitutions that were more common in B35+ individuals than in B35- individuals, but this was one of the six that had no B35 associated pattern of mutation.

**HXB2 Location** RT (432–441)

**Author Location** Pol (587–596)

Epitope EPIVGAETFY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35, B51)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot

granzyme B

Keywords characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- None of seven patients responded to this peptide with GzB producing cells, while two of the patients responded with IFN-gamma producing cells.

**HXB2 Location** RT (434–447)

**Author Location RT (LAI)** 

**Epitope IVGAETFYVDGAAS** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*6802)

**References** Menendez-Arias *et al.* 1998; van der Burg *et al.* 1997

- Recognized by CTL from a long-term survivor that recognized a set of 5 overlapping peptides spanning IVGAETFYVDGAAS as well as PIVLPEKDSW and KITTESIVIW.
- A\*6802 is a subset of HLA-A28.
- This epitope spans the Pol p66 RT p15 (RNAse) domain.

**HXB2 Location** RT (436–445)

Author Location (C consensus)

**Epitope** GAETFYVDGA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6802)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- GAETFYVDGA is an optimal epitope.

**HXB2 Location** RT (436-445)

Author Location RT (436-445)

Epitope GAETFYVDGA

Immunogen

Species (MHC) human (A\*6802)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an A\*6802 epitope.

**HXB2 Location** RT (436–445)

**Author Location** RT (591–600 IIIB)

**Epitope** GAETFYVDGA

Immunogen HIV-1 infection

Species (MHC) human (B45)

References Menendez-Arias et al. 1998

• This epitope spans the Pol p66 RT – p15 (RNAse) domain.

**HXB2 Location** RT (436–445)

Author Location Pol (591–600 IIIB)

Epitope GVETFYVDGA

Immunogen HIV-1 infection

Species (MHC) human (B45)

**Keywords** responses in children, mother-to-infant transmission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- No variants of this epitope were found in a non-transmitting mother who had a CTL response to it.
- This epitope spans the Pol p66 RT p15 (RNAse) domain.

**HXB2 Location** RT (436–452)

**Author Location** (C consensus)

Epitope GAETFYVDGAANRETKI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*3402)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

**References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location RT (436–452) Author Location (C consensus)

Epitope GAETFYVDGAANRETKI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (437–445)

**Author Location** 

Epitope AETFYVDGA

**Epitope name** Pol-AA9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*4501)

**Donor MHC** A\*3002 A\*3201 B\*4501 B\*5301 Cw\*0401

Cw\*1202

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- Subject 00RCH33 was on HAART had a viral load of 2900 and CD4 count of 727 and also recognized the epitopes YPLTFG-WCY, Nef(135-143), HLA B\*5301; RSLYNTVATLY, p17(76-86), HLA A\*3002; and HIGPGRAFY, gp160(310-318), HLA A\*3002.
- Among HIV+ individuals who carried HLA B45, 3/9 (33%) recognized this epitope.

**HXB2 Location** RT (437–447)

**Author Location** RT (592–602 LAI)

**Epitope** AETFYVDGAAN

Subtype B

Immunogen

Species (MHC) human (A28)

**References** Brander & Walker 1996; Menendez-Arias *et al.* 1998

- P. Johnson, pers. comm.
- This epitope spans the Pol p66 RT p15 (RNAse) domain.

**HXB2 Location** RT (437–447)

Author Location Pol (592–602)

Epitope AETFYVDGAAN

Immunogen HIV-1 infection

Species (MHC) human (A28)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** RT (438–448)

**Author Location** (C consensus)

**Epitope** ETFYVDGAANR

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*66)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- ETFYVDGAANR is an optimal epitope.

**HXB2 Location** RT (438–448)

**Author Location** RT (593–603 IIIB)

**Epitope** ETFYVDGAANR

Immunogen HIV-1 infection

Species (MHC) human (A26)

References Menendez-Arias et al. 1998

• This epitope spans the Pol p66 RT – p15 (RNAse) domain.

**HXB2 Location** RT (438–448)

Author Location Pol (593–603 IIIB)

Epitope ETFYVDGAANR

Immunogen HIV-1 infection

Species (MHC) human (A26)

**Keywords** responses in children, mother-to-infant transmission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- One other variant was found that gave a positive, though reduced, CTL response: ETYYVNGAANR.
- This epitope spans the Pol p66 RT p15 (RNAse) domain.

**HXB2 Location** RT (438–448)

**Author Location** 

**Epitope** ETFYVDGAANR

Epitope name ER11

Immunogen

Species (MHC) human (A66)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a A66 epitope.

**HXB2 Location** RT (440–448)

**Author Location** Pol (594–602 SF2)

Epitope FYVDGAANR

Subtype B

Immunogen HIV-1 infection, computer prediction

Species (MHC) human (A\*3303)

Assay type Chromium-release assay

Keywords binding affinity, computational epitope predic-

tion

References Hossain et al. 2003

- HLA-A\*3303 is a common HLA allele in east and southeast Asia. Pol, Gag and Nef SF2 proteins were scanned for potential A\*3303 epitopes. 99 potential epitopes were synthesized, and 52/99 bound to A\*3303. Six of these served as peptide-targets for lysis by PBMC from infected individual, and clones derived from 4 of these 6 could lyse HIV-vaccinia infected target cells, indicating proper processing.
- This epitope is one of the 4 that are properly processed.

**HXB2 Location** RT (448–457)

**Author Location RT** 

Epitope RETKLGKAGY

Immunogen HIV-1 infection

Species (MHC) human (A29)

**Keywords** rate of progression

References van der Burg et al. 1997

- Patients studied were from the Amsterdam cohort.
- CTL epitopes of 3 rapid progressors were compared to 4 longterm survivors (LTS) and no differences could be found in the degree of conservation between them.
- Epitope recognized by a LTS.
- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT.

**HXB2 Location** RT (449–457)

**Author Location** 

**Epitope** ETKLGKAGY

**Epitope name** Pol-EY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2601)

**Donor MHC** A\*3303 A\*2601 B\*5801 B\*8201 Cw\*0302

Cw\*0701

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- Subject 03RCH40 was African American, had a viral load of 2500, CD4 count of 372, was not on HAART, and also recognized the epitope DILDLWIY, Nef(108-115), HLA Cw\*0701.
- Among HIV+ individuals who carried HLA A26, 2/8 (25%) recognized this epitope.

**HXB2 Location** RT (449–457)

**Author Location** Pol (604–612)

Epitope ETKLGKAGY

Immunogen HIV-1 infection

Species (MHC) human (A\*2601)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** RT (449–457)

**Author Location** Pol (604–612)

Epitope ETKLGKAGY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2601)

Country Japan.

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay, Other, HLA binding

**Keywords** immunodominance, characterizing CD8+ T

cells, optimal epitope

References Satoh et al. 2005

- Reverse immunogenetics was used to identify HIV-1 epitopes presented by HLA-A\*2601. Four epitopes endogenously presented by this allele induced peptide-specific CD8 T-cells. HIVinfected individuals predominantly detected 2 of the epitopes, which might be useful for vaccine development. HLA-A\*2601 is common in Asia.
- Immunodominant epitope recognized in 4/6 HIV-infected individuals with HLA-A\*2601. This epitope is highly conserved in clade E (CRF01), and moderately conserved in clade B.

**HXB2 Location** RT (451–459)

**Author Location** Pol (606–)

Epitope KLGKAGYVT

Epitope name Pol606

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: RT Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (MHC) human (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was an intermediate A2 binder that did not induce CTL or CD8+ T-cell IFN gamma responses in transgenic mice. Responses were detected in 2/17 HIV+ HLA-A2 subjects.

**HXB2 Location** RT (467–484)

**Author Location** (C consensus)

Epitope VSLTETTNQKTELQAIQL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*18)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (481-505)

**Author Location** RT (648–672 PV22)

Epitope AIYLALQDSGLEVNIVTDSQYALGI

Immunogen HIV-1 infection Species (MHC) human (B14)

> References Kalams et al. 1994; Menendez-Arias et al. 1998

- · A CTL response used to study gene usage in HLA-B14 re-
- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT.

**HXB2 Location** RT (481–505)

**Author Location** RT (648–672)

Epitope AIYLALQDSGLEVNIVTDSQYALGI

Immunogen HIV-1 infection

Species (MHC) human

References Menendez-Arias et al. 1998; Price et al. 1995

- Study of cytokines released by HIV-1 specific activated CTL.
- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT.

**HXB2 Location** RT (485–493)

**Author Location** RT (640–648 HXB2R)

Epitope ALQDSGLEV Immunogen HIV-1 infection Species (MHC) human (A\*0201)

References Brander et al. 1995; Brander et al. 1996

- This epitope was recognized by PBMC from 3/14 HIV+ asymptomatic patients.
- · This epitope was used along with Env CTL epitope TLTSC-NTSV and a tetanus toxin T helper epitope for a synthetic
- This vaccine failed to induce a CTL response, although a helper response was evident.
- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT.

**HXB2 Location** RT (485–493)

**Author Location** RT (640–648 HXB2R)

Epitope ALQDSGLEV

Immunogen vaccine

Strain: B clade HXB2 HIV component: RT

Species (MHC) human (A2)

References Brander et al. 1995

- Epitope studied in the context of inclusion in a synthetic vac-
- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT.

**HXB2 Location** RT (485-493)

Author Location Pol (649-659 BH10, LAI)

**Epitope** ALQDSGLEV Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

· This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.

• A comprehensive analysis of 160 class I T cell responses in 578 • This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is IYLALQDS-GLE) has similarity with the epidermal growth factor receptor kinase substrate EPS8, fragment ISAAASDSGVE.

**HXB2 Location** RT (485–494)

Author Location RT (485–495 HXB2)

Epitope ALQDSGSEVN

Epitope name 51H Subtype B Immunogen vaccine

> Vector/Type: DNA Strain: multiple epitope immunogen HIV component: p17/p24 Gag,

Pol Adjuvant: IL-12

**Species (MHC)** transgenic mouse (A2)

Assay type Cytokine production, CD8 T-cell Elispot -

IFNγ, Chromium-release assay

Keywords vaccine-specific epitope characteristics, vaccine antigen design

References Bolesta et al. 2005

- · Immunization of transgenic mice with a codon-optimized hGagp17p24-Polp51 DNA plasmid, consisting of clusters of highly conserved CTL epitopes presented by multiple MHC class I alleles, induced 2- to 5-fold higher CD8+ T-cell responses than the corresponding full-length proteins. The modified proteins had the ribosomal frameshift deleted, as well as the potentially immunosuppressive p15, and protease and integrase. This correlated with higher protection against challenge with Gag and Pol expressing recombinant vaccinia virus. Mice immunized with the hGagp17p24-Polp51 also showed an elevated level of type 1 cytokine production as well as an increased titer of p24- and RT-specific IgG2 antibody responses.
- This was 1 of 4 A2 gag/pol epitopes tested. Transgenic mice immunized with the deleted construct induced more potent EliSpot reactions to this epitope than those immunized with full length Gag/Pol.

**HXB2 Location** RT (485–505)

Author Location RT (648-672)

Epitope ALQDSGLEVVTDSQYALGI

Immunogen HIV-1 infection Species (MHC) human (B14)

References Brander & Walker 1995

- Unpublished, S. Kalams.
- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT.

**HXB2 Location** RT (492-501)

**Author Location** Pol (647–656)

**Epitope** EVNIVTDSQY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2601)

Country Japan.

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay, Other, HLA binding

Keywords immunodominance, optimal epitope

References Satoh et al. 2005

- Reverse immunogenetics was used to identify HIV-1 epitopes presented by HLA-A\*2601. 110 peptides were predicted to bind to HLA-A\*2601. 24 of these were demonstrated to bind through a HLA-A\*2601 stabilization assay. Four of these, including this one, were shown to be epitopes endogenously presented by this allele, that can induce peptide-specific CD8 T-cells. HLA-A\*2601 is common in Asia.
- This epitope was recognized in only 1/7 HLA-A\*2601 HIV infected individuals.

**HXB2 Location** RT (492–506)

**Author Location** (C consensus)

Epitope EVNIVTDSQYALGII

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*3910)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (492–506)

**Author Location** (C consensus)

Epitope EVNIVTDSQYALGII

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0802)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (495-503)

**Author Location** 

Epitope IVTDSQYAL

Epitope name IL9

Immunogen

Species (MHC) human (Cw\*0802)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a Cw\*0802 epitope.

**HXB2 Location** RT (496–505)

**Author Location** 

Epitope VTDSQYALGI

Epitope name Pol-VI10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*1503)

Donor MHC A\*3002 A\*6801 B\*0801 B\*1503 Cw\*0701 Cw\*08

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- Subject 01RCH51 was an African American on HAART, viral load 980, CD4 count 811.
- Among HIV+ individuals who carried HLA B15, 1/17 (6%) recognized this epitope.

**HXB2 Location** RT (496–505)

Author Location Pol (651–660)

Epitope VTDSQYALGI

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location RT (496-505)

Author Location Pol (651–660)

Epitope VTDSQYALGI

Epitope name VI10 Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Donor MHC** A\*02, A\*68, B\*14, B\*52, Cw\*08, Cw\*12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** escape, optimal epitope

References Koibuchi et al. 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- The B consensus form of this epitope, VTDSQYALGI, persisted throughout 6 years of chronic infection in 1 individual.

**HXB2 Location** RT (496–505)

**Author Location** Pol (subtype B)

Epitope VTDSQYALGI

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B14, B\*1402)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

• HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi - these CTL may confer protection.

- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found - B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among A, B and D clade viruses.

**HXB2 Location** RT (496-505) **Author Location** RT (663–672 IIIB) **Epitope** VTDSQYALGI Immunogen HIV-1 infection Species (MHC) human (Cw8)

References Brander & Walker 1996

- Unpublished, P. Johnson.
- Published in this database in 1995 as B14, but B14 transfected cells did not present the peptide and it is thought to be presented by the genetically linked Cw8 molecule instead Brander & Walker [1996]

**HXB2 Location** RT (496–505)

**Author Location RT** 

**Epitope** VTDSQYALGI

**Immunogen** HIV-1 exposed seronegative

Species (MHC) human (Cw8)

**Keywords** subtype comparisons, HIV exposed persis-

tently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades - such crossreactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A and D subtype consensus are identical to the B clade
- Thought to be HLA-Cw8 restricted, not B14 as originally reported (C. Brander, B. Walker, and S. Rowland-Jones, personal communication)
- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT.

**HXB2 Location** RT (497–512) Author Location (C consensus)

Epitope TDSQYALGIIQAQPDK

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*0205) Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** RT (502–517) **Author Location** (C consensus) Epitope ALGIIQAQPDKSESEL Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*3901)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression

References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location RT (509-518)

**Author Location** Pol

**Epitope QPDKSESELV** 

Immunogen

Species (MHC) human (B7)

References De Groot et al. 2001

- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT. The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.
  - A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes were identified that could stimulate IFN $\gamma$  production in an ELISPOT assay.
  - QPDKSESELV was newly identified as an HLA-B7 epitope in this study.

**HXB2 Location** RT (509-518)

**Author Location** Pol

Epitope OPDKSESELV

Epitope name 1302

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A03, A24, B07, B38, Cw07, Cw12/13

Country United States. Assay type T-cell Elispot

Keywords binding affinity, computational epitope prediction

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for QPDKSESELV: 36%

**HXB2 Location** RT (516-525)

Author Location RT (516-525)

Epitope ELVNQIIEQL

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Haas et al. 1998

• Of 98 patients in cross-sectional analysis, 78% had CTL against pol - RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)

- New clusters of epitopes were defined utilizing different HLA molecules.
- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT.

HXB2 Location RT (520–528) Author Location Pol (520–528 LAI) Epitope QIIEQLIKK

Subtype B

Immunogen

**Species (MHC)** human (A\*1101) **Keywords** optimal epitope

References Frahm et al. 2007; Fukada et al. 1999

• C. Brander notes this is an A\*1101 epitope.

HXB2 Location RT (520–528)

Author Location Pol (675–683)

Epitope QIIEQLIKK

Subtype B, CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A\*1101)

Keywords subtype comparisons, TCR usage

References Fukada et al. 2002

- Counterparts for eight known clade B HLA A\*1101 epitopes were generated for clade E (CRF01). Three epitopes, identical among clade A-E, were cross-reactive and recognized by clade E infected individuals. The clade E and B analogs to three more HLA A\*1101 epitopes was recognized in a clade specific manner. Two other HLA A\*1101 clade B defined epitopes were found not to have stimulated a response in clade E infected individuals.
- QIIEQLIKK was found to elicit clade-specific responses in clade B (QIIEQLIKK is most common) and clade E (qiieElikk is most common). QIIEQLIKK was strongly recognized by CTL from 1/5 B clade infected Japanese subjects, and qiieElikk from 3/7 E clade infected Thai subjects. The variant qiieKliEk, common in the A subtype, was also recognized in 2/7 E clade infected Thai subjects.
- The binding of QIIEQLIKK, qiieElikk and qiieKliEk to HLA A\*1101 was similar, but CTL clones from individuals did not cross-react with the cross-clade peptides indicating that the substitutions inhibited TCR interaction.

HXB2 Location RT (520–528)
Author Location RT (80–88)
Epitope QIIEQLIKK
Immunogen HIV-1 infection
Species (MHC) human (A\*1101)
Keywords optimal epitope
References Frahm et al. 2007

HXB2 Location RT (520–528)
Author Location Pol (676–684)
Epitope QIIEQLIKK
Epitope name QK9
Subtype B

Immunogen HIV-1 infection **Species (MHC)** human (A11)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

**Keywords** optimal epitope **References** Allen *et al.* 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- This epitope did not vary.

HXB2 Location RT (520–528)

Author Location Pol (676–684)

Epitope QIIEQLIKK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$  References Allen *et al.* 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** RT (520-528)

**Author Location** Pol

Epitope QIIEQLIKK

Epitope name 1336 Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A03, A23, B49, B57

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope prediction

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for QIIEQLIKK: 48%

HXB2 Location RT (530–538)
Author Location
Epitope KVYLAWVPA
Epitope name Pol-KA9
Subtype B
Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Donor MHC A\*0202 A\*0301 B\*4501 B\*5301 Cw\*0401

Cw\*1502

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a
- This epitope was newly defined in this study.
- Patient 04RCH86 was Hispanic, not on HAART, and had a viral load of 7600 and CD4 count of i774.
- Among HIV+ individuals who carried HLA A\*03, 2/21 (10%) recognized this epitope.

**HXB2 Location** RT (530–538)

Author Location Pol (685–693)

Epitope KVYLAWVPA

Epitope name KA9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, Chromium-release assay

Keywords escape, acute/early infection

References Bansal et al. 2005

- Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- The response to this epitope was barely detectable until month

**HXB2 Location** RT (530–538)

Author Location Pol (680–691 BH10, LAI)

Epitope KVYLAWVPA Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is IKKEKVY-LAWV) has similarity with B-cell growth factor precursor, fragment IKKERLWLGPV.

**HXB2 Location** RT (530-540)

Author Location (C consensus)

Epitope RVYLSWVPAHK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords epitope processing, rate of progression, opti-

mal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in a residue outside of the optimized epitope of RVYLSWVPAHK are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** RT (532-540)

**Author Location** Pol (687–)

**Epitope** YLAWVPAHK

Epitope name Pol687

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: RT Adjuvant: Incomplete Freund's Adjuvant

(IFA)

**Species (MHC)** human, transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

- References Corbet et al. 2003 • HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was an intermediate A2 binder, and induced a CD8+ T-cell IFN gamma response in 1/6 mice. Responses were detected in 1/17 HIV+ HLA-A2 subjects.

**HXB2 Location** RT (532–540)

**Author Location** Pol (714–722)

**Epitope** YLAWVPAHK

Immunogen HIV-1 infection

**Species (MHC)** human (A3 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 5/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

Author Location RT (532–540)

Author Location RT (532–540)

Epitope YLAWVPAHK

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Haas et al. 1998

- Of 98 patients in cross-sectional analysis, 78% had CTL against pol – RT was more immunogenic than Integrase and Protease (81%, 51%, and 24% of 37 patients, respectively)
- New clusters of epitopes were defined utilizing different HLA molecules.
- This epitope occurs in the p15 (RNAse) domain of Pol p66 RT.

HXB2 Location RT (532–540)

Author Location RT Pol (687–695)

Epitope YLAWVPAHK

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 3/7 patients recognized this epitope.

## II-B-13 RT-Integrase CTL/CD8 + epitopes

**HXB2 Location** RT-Integrase (553–2)

**Author Location** Pol

**Epitope** STGIRRVLFL

Epitope name SL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- An escape mutation at position 6 (stgirKvlfl) was found not to correspond to the most polymorphic residues in the epitope.
   This is a novel partially mapped epitope.

HXB2 Location RT-Integrase (560-8)

**Author Location** Pol (715–723)

Epitope LFLDGIDKA

Immunogen

Species (MHC) human (B81) Keywords optimal epitope References Frahm *et al.* 2007

## II-B-14 Integrase CTL/CD8+ epitopes

**HXB2 Location** Integrase (9–19) **Author Location** (C consensus)

**Epitope** QEEHEKYHSNW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4403)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the E2 and E3 residues of QEEHEKYH-SNW are associated with the presence of the HLA presenting molecule in the host.
- · QEEHEKYHSNW not optimized.

HXB2 Location Integrase (9–19)

**Author Location** Pol

**Epitope** QEEHEKYHSNW

Epitope name QW11

**Subtype** B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 2, qAehekyhsnw, was found not to correspond to the most polymorphic residues in the epitope. This is a novel partially mapped epitope.

HXB2 Location Integrase (20–28)

**Author Location** Pol (762–770)

Epitope RAMASDFNL

Immunogen HIV-1 infection

Species (MHC) human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind three of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

**HXB2 Location** Integrase (22–31)

Author Location Pol (764-773)

Epitope MASDFNLPPV

Immunogen HIV-1 infection

**Species (MHC)** human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind all five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802)

**HXB2 Location** Integrase (28–36)

**Author Location** (C consensus)

Epitope LPPIVAKEI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0705)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- LPPIVAKEI is an optimal epitope for B\*4201, B\*0705, and B\*5101.

HXB2 Location Integrase (28–36)

**Author Location** (C consensus)

Epitope LPPIVAKEI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location Integrase (28–36)

Author Location (C consensus)

Epitope LPPIVAKEI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the P3, I4, A6, and K7 residues of LPPI-VAKEI are associated with the presence of the HLA presenting molecule in the host.
- LPPIVAKEI is an optimal epitope for B\*4201, B\*0705, and B\*5101

**HXB2 Location** Integrase (28–36)

Author Location Pol (743–751 SF2)

Epitope LPPVVAKEI

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

**Keywords** subtype comparisons, rate of progression

References Tomiyama et al. 1999

- HLA-B27, -B51, and -B57 are associated with slow progression to AIDS, while HLA-B35, -B8, -B24 are associated with a rapid progression to AIDS (Nat. Med. 2:405, 1996;Lancet 22:1187, 1986;Hum Immunol 22:73, 1988;Hum Immunol 44:156, 1995)
- 15% of Japanese populations carry HLA-B51 while HLA-B27 and -B57 are detected in less than 0.3%
- Of the 172 HIV-1 peptides with HLA-B\*5101 anchor residues, 33 bound to HLA-B\*5101, seven of these peptides were reactive with CTL from 3 B\*5101 positive individuals, and six were properly processed.
- Four of the six epitopes were highly conserved among B subtype sequences – LPPVVAKEI is highly conserved.

HXB2 Location Integrase (28–36)

**Author Location** (C consensus)

Epitope LPPIVAKEI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- LPPIVAKEI is an optimal epitope for B\*4201, B\*0705, and B\*5101.

HXB2 Location Integrase (28–36)

**Author Location** Integrase (28–36 HXB2)

Epitope LPPVVAKEI

Epitope name LI9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

**Donor MHC** A\*0201, A\*2501,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

• T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.

HXB2 Location Integrase (28–36)

**Author Location** 

**Epitope** LPPIVAKEI

Epitope name LI9

Immunogen

Species (MHC) human (B42)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B42 epitope.

HXB2 Location Integrase (28–36)

**Author Location** Pol (28–36)

Epitope LPPVVAKEI

Immunogen HIV-1 infection

Species (MHC) human (B51)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DO7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

**Keywords** rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 7 epitopes from this individual that varied over time, although the internal mutations did not become fixed. The lppIvakei variant arose at intermediate time points.

HXB2 Location Integrase (28–36)

Author Location Integrase (28-36)

Epitope LPPIVAKEI

Epitope name LI9

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human (B51)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

assay, HLA binding

Keywords subtype comparisons, computational epitope

prediction, mother-to-infant transmission, escape, reversion, viral fitness, optimal epitope

References Leslie et al. 2005

B\*1801, B\*5101, • An I4V substitution (LPPvVAKEI) is suggested to be driven by CTL escape in B51-positive subjects. The escape form is the consensus form of the epitope in the B clade, and stable in the absence of HLA-B51. In the C clade the B51 is rare, and the Val escape mutation is also rare.

**HXB2 Location** Integrase (29–46)

**Author Location** (C consensus)

Epitope PPIVAKEIVASCDKCQLK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*8101)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Integrase (62–71)

**Author Location** Pol

**Epitope QLDCTHLEGK** 

Epitope name 1335

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A03, A23, B49, B57; A03, A11, B14, B05,

Cw08

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for OLDCTHLEGK: 61%.

HXB2 Location Integrase (66–74) Author Location (C consensus) Epitope THLEGKIIL

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*1510)

**Country** South Africa. **Assay type** CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the L9 residue of THLEGKIIL are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** Integrase (66–74)

**Author Location** 

Epitope THLEGKIIL

Epitope name TIL9

Immunogen

Species (MHC) human (B\*1510)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*1510 epitope.

**HXB2 Location** Integrase (82–89) **Author Location** RT (797–804 SF2)

Epitope GYIEAEVI Immunogen HIV-1 infection Species (MHC) human (A\*2402) References Ikeda-Moore *et al.* 1997

- Defined using reverse immunogenetics 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) 53 of the 59 peptides bound A\*2402.
- This peptide induced CTL in 1/4 HIV-1 + people tested.
- GYIEAEVI bound to A\*2402 weakly, the epitope can be processed in a vaccinia construct and presented two specific CTL clones were obtained.

HXB2 Location Integrase (89–98)

**Author Location** Pol

Epitope IPAETGQETA

Immunogen

Species (MHC) human (B56)

References De Groot et al. 2001

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 semight serve as epitopes.
   The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.
  - A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes could stimulate IFNγ production in an ELISPOT assay.
  - IPAETGQETA was newly identified as an HLA-B56 epitope in this study.

HXB2 Location Integrase (89–98)

**Author Location** Pol

Epitope IPAETGQETA

Epitope name 1294

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A02, A03, B07, B58, Cw07

Country United States.
Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for IPAETGQETA: 8%

**HXB2 Location** Integrase (89–98)

Author Location Pol (805-814 BH10, LAI)

Epitope IPAETGQETA Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is PAETGQETAY) has similarity with Integrin beta-4 precursor (GP150)(CD104), fragment PAETNGEITAY.

**HXB2 Location** Integrase (90–107)

Author Location (C consensus)

Epitope PAETGQETAYFILKLAGR

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6802)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location Integrase (92–99)

**Author Location** (C consensus)

**Epitope** ETGQETAY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2601)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- · ETGQETAY is an optimal epitope.

**HXB2 Location** Integrase (96–104)

**Author Location** Integrase (823–831)

Epitope ETAYFILKL

Immunogen

Species (MHC) human (A\*6802)

Keywords subtype comparisons

References Dong & Rowland-Jones 1998

 Epitope found in clade A, B, and D – pers. comm. S. Rowland-Jones and T. Dong.

HXB2 Location Integrase (96–104)

**Author Location** Pol (subtype A)

**Epitope** ETAYFILKL

Subtype A

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A\*6802)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix systemic CD8+ T cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

HXB2 Location Integrase (96–104)

**Author Location** Pol

Epitope ETAYFILKL

Immunogen HIV-1 infection

Species (MHC) human (A\*6802)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted – 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.

- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls (ML1671)

HXB2 Location Integrase (96–104)

**Author Location** Pol (744–752)

Epitope ETAYFILKL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A\*6802)

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

- ETAYFILKL cross-reacts with clades A, B and D.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-A\*6802 women, 3/12 HEPS and 9/11 HIV-1 infected women recognized this epitope likelihood ratio 7.9, p value 0.01, and HEPS women tended to respond to DTVLEDINL, while infected women to ETAYFILKL.
- The dominant response to this HLA allele was to this epitope in 2 of the 3/12 HEPS cases and in all 9/11 HIV-1 infected women that responded to the epitope.
- Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.
- Subject ML 1203 started with CTL responses to A\*6802 DTVLEDINL and to B7 FPVTPQVPLR prior to seroconversion, and upon seroconversion acquired additional responses to A\*6802 ETAYFILKL which became dominant, B7 TPGPG(V/I)RYPL, B7 IPRRIRQGL, and B7 SPRTLNAWV.
- Subject ML 1707 started with a CTL response to A\*6802 DTVLEDINL prior to seroconversion, and switched to A\*6802 ETAYFILKL and A24 RDYVDRFFKTL post-seroconversion.
- Subject ML 1830 made no detectable response prior to seroconversion, but responded to A\*6802 DTVLEDINL and A\*6802 ETAYFILKL post-seroconversion.

HXB2 Location Integrase (96–104)

**Author Location** Pol (744–752)

**Epitope** ETAYFILKL

Immunogen HIV-1 infection

Species (MHC) human (A\*6802)

References Appay et al. 2000

• This epitope is newly defined in this study.

- · Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virusspecific CD8+ T cells produced IFN- $\gamma$  and MIP-1 $\beta$  with a distinct subset that failed to produce TNF- $\alpha$

HXB2 Location Integrase (101–111)

**Author Location** (C consensus)

**Epitope** ILKLAGRWPVK

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*0301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords epitope processing, rate of progression, opti-

mal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in a residue outside of the optimized epitope of ILKLAGRWPVK are associated with the presence of the HLA presenting molecule in the host.

HXB2 Location Integrase (105–121)

**Author Location** (C consensus)

Epitope AGRWPVKVIHTDNGSNF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Integrase (123–132)

**Author Location** Integrase (123–132)

Epitope STTVKAACWW

Immunogen

Species (MHC) human (B57)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** Integrase (123–132)

**Author Location** Integrase

Epitope STTVKAACWW

**Epitope name** SW10

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Chromium-release assay, Flow

cytometric T-cell cytokine assay

Keywords epitope processing, supervised treatment in-

terruptions (STI), rate of progression, immun-

odominance

References Rodriguez et al. 2004

- Protease and integrase are shown to be frequently targeted by CD8 T-cell responses (23% and 68% of 56 HIV+ patients, respectively). Responses tend to cluster in conserved regions of Int, although 1 high conserved region had no responses. CTL frequencies per unit protein length for Pro and Int were similar to other HIV non-structural proteins. Three novel HLA class I-restricted optimal epitopes were found and characterized with fine mapping.
- All 5 HLA-B57 patients recognized this epitope and were longterm nonprogressors.

HXB2 Location Integrase (123-132)

**Author Location** Pol

**Epitope STTVKAACWW** 

**Epitope name** SW10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Two escape mutations, at positions 2 and 3, were found in the most polymorphic residue in the epitope. These were shared between clades B and C. The T840N mutation at residue 2, sNtvkaacww, was significantly more common in persons expressing HLA-B57, often in conjunction with T841A or V sT[A/V]kaacww.

HXB2 Location Integrase (123–132)

**Author Location** 

**Epitope** STTVKAACWW

Immunogen HIV-1 infection

Species (MHC) human (B57, B\*5801)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords responses in children, mother-to-infant transmission, characterizing CD8+ T cells

References Feeney et al. 2005

• Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.

HXB2 Location Integrase (127–135)
Author Location Pol (869–877)
Epitope KAACWWAGI
Immunogen HIV-1 infection
Species (MHC) human (A2 supertype)
Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind three of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

**HXB2 Location** Integrase (135–143)

**Author Location** (C consensus)

Epitope IQQEFGIPY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the Q2 residue of IQQEFGIPY are associated with the presence of the HLA presenting molecule in the host.

HXB2 Location Integrase (135–143)

**Author Location** 

Epitope IQQEFGIPY

Immunogen

Species (MHC) human (B\*1503)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B\*1503 epitope.

HXB2 Location Integrase (135–143)

**Author Location** Integrase (135–143)

Epitope IQQEFGIPY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Assay type Other

**Keywords** HLA associated polymorphism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- IQQEFGIPY was a previously defined B\*1503 presented epitope that encompassed a polymorphism, IqQEFGIPY,in the second position.

HXB2 Location Integrase (157–166)

**Author Location** (C consensus)

Epitope ELKKIIGQVR

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*33)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- ELKKIIGQVR is an optimal epitope.

HXB2 Location Integrase (164–172)

Author Location (C consensus)

Epitope QVRDQAEHL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0205)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- QVRDQAEHL is an optimal epitope.

**HXB2 Location** Integrase (165–172)

Author Location Integrase (165–172)

Epitope VRDQAEHL

Epitope name VL8

Immunogen

Species (MHC) human (Cw\*18)

**Keywords** optimal epitope References Frahm et al. 2007

• C. Brander notes this is a Cw18 epitope.

HXB2 Location Integrase (165–172)

Author Location (C consensus)

Epitope VRDQAEHL Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- VRDQAEHL is an optimal epitope.

**HXB2 Location** Integrase (165–172)

**Author Location** Integrase (165–172)

**Epitope** VRDQAEHL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1801)

Assay type Other

**Keywords** HLA associated polymorphism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- VRDQAEHL was a previously defined Cw1801 presented epitope that encompassed a Cw18 associated polymorphism, VRdQAEHL,in the third position.

HXB2 Location Integrase (171–180)

**Author Location** Pol

Epitope HLKTAVQMAV

Epitope name 1247

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A01, A02, B08, Cw16

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC
- Estimated binding probability for HLKTAVOMAV: 82%

HXB2 Location Integrase (173-181)

Author Location Pol (888–896)

Epitope KTAVQMAVF

Immunogen

Species (MHC) human (B\*5701)

Keywords optimal epitope

References Frahm et al. 2007

- C. Brander notes this is a B\*5701 epitope.
- Epitope is motif based, personal communication from C. Hay.
- Subtype of B57 not determined.

**HXB2 Location** Integrase (173–181)

Author Location Pol (888–896)

Epitope KTAVQMAVF

Immunogen

Species (MHC) human (B57)

References Hay 1999

• Epitope is motif based, personal communication from C. Hay.

HXB2 Location Integrase (173–181)

Author Location Pol

Epitope KTAVQMAVF

Epitope name KF9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was quite conserved in people carrying B57, but two substitutions were found in 11 B57+ individuals tested: Rtavqmavf and ktavqmavL.

HXB2 Location Integrase (173–181)

**Author Location** Pol (889–897)

Epitope KTAVQMAVF

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B57)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- ing the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

HXB2 Location Integrase (173–181)

**Author Location** 

Epitope KTAVQMAVF Immunogen HIV-1 infection Species (MHC) human (B57, B\*5801)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords responses in children, mother-to-infant trans-

mission, escape

References Feeney et al. 2005

• Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.

HXB2 Location Integrase (177–186)

Author Location Pol (919–928)

**Epitope** QMAVFIHNFK

Immunogen HIV-1 infection Species (MHC) human (A3 supertype)

**Keywords** supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 5/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location Integrase (178–186)

**Author Location** Pol (920–928)

Epitope MAVFIHNFK

Immunogen HIV-1 infection

**Species (MHC)** human (A3 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Two-thirds of all mutations arising in nonenvelope proteins dur-CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
  - Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
  - A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
  - This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location Integrase (179–187)

**Author Location** Pol (921–929)

Epitope AVFIHNFKR

Immunogen HIV-1 infection

Species (MHC) human (A3 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer
- · Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 5/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

**HXB2 Location** Integrase (179–188)

Author Location Integrase (179–188)

Epitope AVFIHNFKRK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Integrase (179–188)

Author Location Integrase (179–188 LAI)

Epitope AVFIHNFKRK

Subtype B

**Immunogen** 

Species (MHC) human (A\*1101)

Keywords optimal epitope

References Frahm et al. 2007; Fukada et al. 1999

• C. Brander notes this is an A\*1101 epitope.

**HXB2 Location** Integrase (179–188)

Author Location Pol (894–903)

**Epitope** AVFIHNFKRK

Subtype B, CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A\*1101)

**Keywords** subtype comparisons

## References Fukada et al. 2002

- Counterparts for eight known clade B HLA A\*1101 epitopes were generated for clade E (CRF01). Three epitopes, identical among clade A-E, were cross-reactive and recognized by clade E infected individuals. The clade E and B analogs to three more HLA A\*1101 epitopes was recognized in a clade specific manner. Two other HLA A\*1101 clade B defined epitopes were found not to have stimulated a response in clade E infected individuals.
- AVFIHNFKRK is commonly found in viruses representing subtypes A-E. It was strongly recognized by CTL from 4/7 E clade infected Thai subjects.

HXB2 Location Integrase (179–188)

**Author Location** Pol (894–903 93TH253 subtype CRF01)

Epitope AVFIHNFKRK Epitope name P894-903 Subtype CRF01\_AE

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A11)

Keywords HIV exposed persistently seronegative (HEPS)

References Bond et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers - weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not
- This epitope was weakly reactive in the HEPS study subjects 265 who was HLA A2/A11 and 128 who was HLA A11/A33, and had been predicted to be a possible A11 epitope using Epimer in Bond et al. [2001]

HXB2 Location Integrase (179-188)

**Author Location** Pol

Epitope AVFIHNFKRK

Epitope name 1264

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A11, A3, A68)

**Donor MHC** A01, A68, B15, B40, Cw03; A03, A11, B14, B51, Cw08, Cw13; A25, A68, B18, B27

Country United States.

Assav type T-cell Elispot

Keywords binding affinity, supertype, computational epitope prediction, immunodominance, cross-

presentation by different HLA

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC
- Estimated binding probability for AVFIHNFKRK:53% Supertype epitope binding to A11, A03 and A68. Immunodominant.

HXB2 Location Integrase (179–188)

Author Location Integrase (894–904)

**Epitope** AVFIHNFKRK

Epitope name A3-AK10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A3, B7, Cw7

Keywords dynamics, supervised treatment interruptions

(STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 1/7 individuals began to have detectable responses to this epitope after STI.

HXB2 Location Integrase (179–188)

Author Location Integrase (179–188)

Epitope AVFIHNFKRK

Epitope name A3-AK10 Pol

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Assay type CD8 T-cell Elispot - IFNγ

Keywords supervised treatment interruptions (STI), es-

cape, early treatment, superinfection

References Altfeld et al. 2002a

- · An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.
- The second infecting strain had the variant avfVhnfkrk. The CTL response to the second variant was zero at all timepoints. The CTL response to the first variant was low and declined over time.

HXB2 Location Integrase (179–188)

**Author Location** Pol

**Epitope** AVFIHNFKRK

Epitope name 1264

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A03, A23, B49, B57; A03, A24, B27, B57,

Cw13, Cw18

Country United States. Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for AVFIHNFKRK: 52%

**HXB2 Location** Integrase (179–196)

Author Location Pol (894–911)

Epitope AVFIHNFKRKGGIGGYSA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Integrase (185–194)

Author Location Integrase (185–194)

Epitope FKRKGGIGGY

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

**Keywords** optimal epitope

References Frahm et al. 2007

**HXB2 Location** Integrase (185–194)

**Author Location** (C consensus)

Epitope FKRKGGIGGY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location Integrase (185-194)

Author Location (C consensus)

Epitope FKRKGGIGGY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the K4 and G9 residues of FKRKG-GIGGY are associated with the presence of the HLA presenting molecule in the host.

HXB2 Location Integrase (185–194)

**Author Location** Integrase (185–194)

Epitope FKRKGGIGGY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Assay type Other

Keywords HLA associated polymorphism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- FKRKGGIGGY was a previously defined B\*1503 presented epitope that encompassed a polymorphism, FKRkGGIGGY,in the fourth position.

**HXB2 Location** Integrase (210–227)

**Author Location** Pol (925–942)

Epitope TKELQKQIIKIQNFRVYY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Integrase (218–235)

Author Location RT-Integrase (218–235 HXB2)

Epitope TKIQNFRVYYRDSRDPLW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 21% of the study subjects, and it was one of the 25 most frequently recognized peptides.

HXB2 Location Integrase (219–226)

**Author Location** (C consensus)

Epitope KIQNFRYY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the K1 residue of KIQNFRYY are associated with the presence of the HLA presenting molecule in the host.

HXB2 Location Integrase (219–227)

**Author Location** 

Epitope KIQNFRVYY

Epitope name Pol-KY9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*3002)

Donor MHC A\*0205 A\*3002 B\*1402 B\*5301 Cw\*0401

Cw\*0802

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- Patient 00RCH28 was African American, not on HAART, had a viral load of 5900 and CD4 count of 889, and she also recognized RIRQGLERA, gp160(846-854), A\*0205.
- Among HIV+ individuals who carried HLA A30, 6/16 (38%) recognized this epitope.

HXB2 Location Integrase (219-227)

Author Location Integrase (219–227)

Epitope KIQNFRVYY

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Keywords optimal epitope

**References** Frahm *et al.* 2007

HXB2 Location Integrase (219–227)

**Author Location** (C consensus)

Epitope KIQNFRVYY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Country South Africa.

**Assav type** CD8 T-cell Elispot - IFNγ

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location Integrase (219–227)

**Author Location** Integrase

Epitope KIQNFRVYY

Epitope name KY9

Immunogen HIV-1 infection

Species (MHC) human (A30)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Chromium-release assay, Flow

cytometric T-cell cytokine assay

Keywords epitope processing, supervised treatment in-

terruptions (STI), immunodominance

References Rodriguez et al. 2004

 Protease and integrase are shown to be frequently targeted by CD8 T-cell responses (23% and 68% of 56 HIV+ patients, respectively). Responses tend to cluster in conserved regions of Int, although 1 high conserved region had no responses. CTL frequencies per unit protein length for Pro and Int were similar to other HIV non-structural proteins. Three novel HLA class I-restricted optimal epitopes were found and characterized with fine mapping.

**HXB2 Location** Integrase (219–228) **Author Location** Pol (934–943 SF2)

Epitope KIQNFRVYYR

Subtype B

Immunogen HIV-1 infection, computer prediction

Species (MHC) human (A\*3303)

Assay type Chromium-release assay

**Keywords** binding affinity, computational epitope prediction

References Hossain et al. 2003

- HLA-A\*3303 is a common HLA allele in east and southeast Asia. Pol, Gag and Nef SF2 proteins were scanned for potential A\*3303 epitopes. 99 potential epitopes were synthesized, and 52/99 bound to A\*3303. Six of these served as peptide-targets for lysis by PBMC from infected individual, and clones derived from 4 of these 6 could lyse HIV-vaccinia infected target cells, indicating proper processing.
- This epitope is one of the 4 that are properly processed.

**HXB2 Location** Integrase (219–228)

**Author Location** Pol (919–928)

Epitope KIQNFRVYYR
Immunogen HIV-1 infection

Species (MHC) human (A3 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 5/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location Integrase (241–249)

**Author Location** Pol (576–584)

Epitope LLWKGEGAV

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

References van der Burg et al. 1996

- Slow dissociation rate, associated with immunogenicity in transgenic HLA-A\*0201/K<sup>b</sup> mice.
- CTL generated by *in vitro* stimulation of PBMC derived from uninfected individual.

HXB2 Location Integrase (241–249) Author Location RT (956–964 HXB2R)

Epitope LLWKGEGAV

Epitope name LR28 Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant (IFA), Montanide (ISA 720), P30, PLG

Species (MHC) mouse (A\*0201)

**Keywords** binding affinity, vaccine-specific epitope characteristics, immunodominance

References Peter et al. 2001

- The stability of peptide binding to HLA-A2.1 was determined for six HLA-A2.1 peptides included in this vaccine study

   ILKEPVHGV (RT), SLYNTVATL (p17), SLLNATDIAV (gp41) and LLWKGEGAV (RT) all bound with high affinity comparable to a influenza epitope reference (GILGFVFTL), while RGPGRAFVTI and VIYQYMDDL bound with a lower affinity (relative binding activity = 0.01).
- The four high-affinity peptides formed stable complexes with half-lives ranging between 8 and 32 hours, while the low affinity peptides had half lives of less than an hour.
- HLA-A2.1 transgenic mice were immunized with the six HIV-1 peptides and P30, as a universal T-helper epitope, with IFA or Montanide or microspheres as adjuvants.
- All peptides except VIYQYMDDL induced a stong CTL response in Cr-release assays stronger responses were observed when peptides were delivered alone, indicating immunodominance when the combination was used.

**HXB2 Location** Integrase (241–249)

Author Location RT (956–964 HXB2R)

Epitope LLWKGEGAV

Epitope name LR28

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant

(IFA), IL-12, P30

Species (MHC) mouse (A\*0201)

Keywords vaccine-specific epitope characteristics, im-

munodominance

References Peter et al. 2002

• When HIV-1 peptides were used to vaccinate HLA-A2.1 transgenic A2-Kb mice, strong responses to five peptides were observed when the peptides were given individually, but immunodominance limited the response to some of the peptides when they were given in combination Peter *et al.* [2001]. IL-12 can counteract immunodominance in BALB/c mice, so it was given with the multiple epitope vaccination, and was instead found to specifically eliminate the HLA-A2.1-epitope CTL responses, but not Kb CTL responses. This was possibly a consequence of transient depletion of T-cells, B cells and macropahges in the spleen.

HXB2 Location Integrase (241–249)

**Author Location** Pol

Epitope LLWKGEGAV

Epitope name L9V

Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV component: gp140, gp140∆V3

**Species (MHC)** transgenic mouse (A\*0201)

Assay type Chromium-release assay, Flow cytometric Tcell cytokine assay

Keywords memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

· A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted CTL responses against multiple conserved HIV-1 epitopes in HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw virus expressing gp140 with deleted V1, V2, and V3 loops successfully induces neutralizing antibodies against HIV-1. A live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination against HIV and measles.

HXB2 Location Integrase (241–249) Author Location Pol (956–964)

**Epitope** LLWKGEGAV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords dendritic cells

References Kundu et al. 1998b

- · Allogeneic dendritic cells (DCs) were obtained from HLAidentical siblings, pulsed with rgp160 MN or A2-restricted HIV-1 epitope peptides, and infused monthly into six HIVinfected patients.
- · 1/6 showed increased Env-specific CTL and increased lymphoproliferative responses, 2/6 showed increase only in proliferative responses, and 3/6 showed no change – pulsed DCs were well tolerated.
- LLWKGEGAV is a conserved HLA-A2 epitope included in this study – 6/6 patients had this sequence as their HIV direct sequence, but only four of these had a detectable CTL response.

HXB2 Location Integrase (241–249)

Author Location Pol (956-964 HXB2R)

**Epitope** LLWKGEGAV

Immunogen peptide-HLA interaction

Species (MHC) human (A2)

References Parker et al. 1992; Parker et al. 1994

• Studied in the context of HLA-A2 peptide binding.

**HXB2 Location** Integrase (241–249)

Author Location Pol (956–964 HXB2R)

Epitope LLWKGEGAV

Immunogen peptide-HLA interaction

Species (MHC) human (A2)

References Brander et al. 1995

· No CTL activity found in HIV-infected subjects, epitope studied in the context of inclusion in a synthetic vaccine.

**HXB2 Location** Integrase (241–249)

Author Location Pol (956–964)

**Epitope** LLWKGEGAW

Immunogen HIV-1 infection

Species (MHC) human (A2, A\*0201)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location Integrase (260–268)

Author Location Integrase (260-268)

Epitope VPRRKVKII

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*42)

Assay type Other

Keywords epitope processing, HLA associated polymor-

phism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- VPRRKVKII was a previously defined B\*42 presented epitope that was associated with a polymorphism, VPRRKVKIIlk seen just after the last position in that epitope.

HXB2 Location Integrase (260–268)

Author Location Integrase (260–268)

Epitope VPRRKAKII

Immunogen

Species (MHC) human (B42)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** Integrase (263–271)

Author Location Integrase (263–271)

Epitope RKAKIIRDY

Immunogen HIV-1 infection Species (MHC) human (B\*1503)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** Integrase (263–271)

**Author Location** Integrase (263–271)

Epitope RKAKIIRDY

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B\*1503)

**Donor MHC** A\*2301, B\*3501, B\*1503, Cw2, Cw7

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

HXB2 Location Integrase (263–271)

**Author Location** (C consensus)

Epitope RKAKIIKDY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** Integrase (263–271)

**Author Location** (C consensus)

Epitope RKAKIIKDY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords epitope processing, rate of progression, opti-

mal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the A3 residue of RKAKIIKDY are associated with the presence of the HLA presenting molecule in the host. Mutation of a residue outside of the optimized epitope also associated with HLA.

HXB2 Location Integrase (266–275)

Author Location (C consensus)

Epitope KIIKDYGKQM

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the I3 residue of KIIKDYGKQM are associated with the presence of the HLA presenting molecule in the host.
- KIIKDYGKQM not optimized.

## II-B-15 Pol CTL/CD8 + epitopes

HXB2 Location Pol

**Author Location** 

**Epitope** 

Immunogen computer prediction

**Species (MHC)** (A\*0201, B\*3501)

**Keywords** subtype comparisons, computational epitope

prediction

References Schönbach et al. 2002

 Computational methods (artificial neural networks, hidden Markov models, binding matrices based on HLA association rates) were used to identify HLA-A\*0201 and HLA-B\*3501 HIV T-cell epitope candidates from 533 Gag, Env and Pol sequences of which 374 were derived from HIV-1, 97 were derived from HIV-2 and 62 from SIV. Comparisons to known epitopes and between clades were made.

HXB2 Location Pol

Author Location Pol

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human (A\*0201, Cw\*08)

References Shacklett et al. 2000

HIV-1 specific, MHC class I-restricted CTL killing was detected in duodenal and rectal gut associated lymphoid tissue (GALT) sites from three infected individuals – the distribution of class I restricted CTL was different in the peripheral blood samples and GALT samples.

HXB2 Location Pol

**Author Location RT (IIIB)** 

Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords epitope processing, escape

References Moore et al. 2002b

- HIV polymorphisms in the RT protein were examined relation to HLA alleles found in a population of 473 participants in the Western Australian HIV Cohort Study. 64 significant associations between polymorphisms at particular positions and HLA alleles were detected, for HLA-B7, -B12, -B35 and -B15. Fifteen of these were in positions with known epitopes, 4 in anchor residues, 11 in other positions. Six additional polymorphic sites associated with particular HLA molecules flanked known epitopes and may relate to processing.
- 25 negative associations were also found between polymorphism and HLA alleles. The authors propose this is due to escape mutations in epitopes presented by common HLA types dominating in the population, and give examples of five amino acids which are in the consensus and tend to be stable in those with the most common HLA allele, HLA-A2.

HXB2 Location Pol

Author Location Pol

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Assay type** Tetramer binding, Flow cytometric T-cell cytokine assay

Keywords assay standardization/improvement

References Wu et al. 2005

 A flow cytometric assay for validation of HIV-1 gag- or polspecific- CD8/HLA-A2 T-cells was shown to be sensitive and specific, being able to detect HIV-1 CTL at the single T-cell level. An inverse correlation between HIV plasma viremia and gag- and pol-specific-CD8/HLA-A2 T-cells was observed.

HXB2 Location Pol

**Author Location** Pol

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*35)

Keywords rate of progression

References Jin et al. 2002

- Patients with HLA-B\*35 variants B\*3502, B\*3503, B\*3504, and B\*5301 tend to proceed to AIDS more quickly than those with B\*3501.
- Of 32 patients with HLA-B\*35 alleles CD8+ CTL responses were quantified using an intracellular cytokine staining assay – 75% had responses to Pol, 69% to Gag, 50% to Nef, and 41% to Env.
- The overall magnitude of CTL responses did not differ between those bearing B\*3501 and the others. A higher percentage of Gag responses was observed in those that had lower RNA levels that carried B\*3501, and there was a negative association with viral load and CTL activity. The data is consistent with higher levels of CTL responses contributing to protection in B\*3501

individuals, but not in B\*3502, B\*3503, B\*3504, and B\*5301 individuals.

HXB2 Location Pol

**Author Location** Pol

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade HXB2, B clade NL43 HIV component: Gag, Pol

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Huang et al. 2001

- Different HIV strains were used for different regions: gag HXB2, pol NL43
- Mice were immunized with four humanized DNA constructs: GagPol, that would form a pseudoparticle carrying Gag and Pol, Gag, Pol or a GagPol fusion construct.
- The GagPol pseudoparticle, Gag and GagPol fusion construct all elicited strong anti-Gag CTL, but only the GagPol fusion construct elicited strong anti -Pol CTL.

HXB2 Location Pol

**Author Location RT (LAI)** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Buseyne et al. 1998a

 This study showed a correlation between strong CTL memory and breadth of response in 7-12 month old infants, and remaining AIDS-free for the first year of life, higher absolute CD4 and CD8 cells, and lower viral load.

**HXB2** Location Pol

**Author Location** p66 (LAV)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, dendritic cells

References Zheng et al. 1999

- Protein delivery (gp160 LAV, p66 LAV, and p24 NY5) to human dendritic cells (DC) with liposomes provides enhanced memory CTL response relative to delivery of protein alone.
- Chloroquine administration enhanced epitope presentation, and brefeldin A and peptide aldehyde inhibitors inhibited antigen presentation, suggesting epitopes were processed by classical proteasome pathway.

HXB2 Location Pol

Author Location Pol (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression, Th1

References Wasik et al. 2000

- HIV+ infants that progressed rapidly to AIDS had lower Th1 responses and decreased production of beta-chemokines and IL-2 relative to other HIV+ infants.
- No HIV+ infants had no demonstrable CTL at birth, but Th1 responses accompanied by CTL responses developed in children with slowly progressive disease, and not in rapid progressors.

 CTLp frequencies were determined by limiting dilution using autologous B cells infected with vaccina/HIV constructs.

HXB2 Location Pol Author Location Pol (LAI) Epitope Subtype B Immunogen vaccine

Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag, gp41, Protease, V3

Species (MHC) human

References Salmon-Ceron et al. 1999

- The vaccine used was a rec canarypox with HIV-1 gp120 MN, tm/gag/protease LAI (vCP205), alone or with p24E-V3 MN synthetic peptide (CLTB-36))
- Twenty HIV negative subjects were vaccinated in phase I trial with combinations of vCP205 and CLTB-36.
- Immunization with vCP205 induced HIV-1-specific ABs to gp120, V3, and p24 antigens, and CTL immune responses against vCP205 were detected after the fourth immunization in 33% of the subjects against Env, Gag and Pol, but the CLTB-36 peptide did not produce AB or CTL immune responses against p24 or gp160.

HXB2 Location Pol

Author Location Pol (172–219 subtype B)

Epitope Subtype B Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade LAI, B clade SF2 HIV component: Env. Gag. Nef, Protease

Species (MHC) human

References Gorse et al. 1999b

- The vaccine used was rec canarypox expressing HIV-1 env, gag, pol, nef and protease (vCP300) with or without administration of HIV-1 SF-2 rgp120.
- In vitro inducible CTL activity against HIV-1 Env, Gag, Pol, and Nef antigens was observed in 79% (15 of 19) of vaccine recipients.
- The combination of vCP300 and vP1291 together resulted in an overall increase in CTL induction and detection sensitivity.

HXB2 Location Pol

Author Location Pol (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Betts *et al.* 1999

This study demonstrated an inverse correlation between HIV
Type I plasma viral load and CTL activity directed against
HIV-1 Pol, and stronger combined effects of Pol- and Envspecific CTL, in long-term survivors (LTS) of HIV-1 infection.

HXB2 Location Pol

Author Location Pol (BRU)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Aladdin *et al.* 1999

• In vitro measurements of CTL-activity by Cr release assay in bulk culture showed no correlation between CTL-activity (gp120, Gag, Pol and Nef) and disease progression as measured by viral load, CD4 and time to death.

HXB2 Location Pol

Author Location RT (LAI)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Buseyne *et al.* 1998b

 In infants with positive CTL responses, most responses showed cross-clade reactivity with somewhat diminished recognition of epitopes from different subtypes.

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env, Gag, Pol, Vif Adjuvant: B7, IL-12

Species (MHC) mouse

References Kim et al. 1997c

- A gag/pol, vif or gp160 DNA vaccine, when delivered in conjunction with the plasmid encoding the co-stimulatory molecules B7 and IL-12, gave a dramatic increase in both the cytotoxic and proliferative responses in mice.
- When IL-12 was present, CTL response could be detected even without in vitro stimulation.

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Trickett et al. 1998

- Twelve HIV-1 infected patients were re-infused with their own lymphocytes, cryopreserved from an earlier time point in the infection.
- Improvement in CD4+ and CD8+ T cells were seen in 7/12, and an increase in the CTL response to Pol was seen in one patient.

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Froebel *et al.* 1997

 Two HIV-1 infected children with contrasting disease courses were followed longitudinally – one died of AIDS, the other is a long-term non-progressor.

- Reactivity against Gag, Pol, Env and Tat proteins was tested by PBMC bulk cultured cells reacting with protein expressed in vaccinia constructs in autologous EBV transformed B cells.
- The child who progressed consistently had CTL against Pol
- The long-term non-progressing child had no detectable CTL, but was heterozygous for a mutation in the CCR5 receptor and for HLA-B49, which has been shown to be associated with slower progression.

HXB2 Location Pol

Author Location Pol (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons References Betts et al. 1997

- 6/8 individuals from Zambia infected with C clade virus had CTL that were able to make response to B clade HIV-1 IIIB vaccinia-expressed Gag, Pol and Env proteins.
- A vigorous cross-clade response was not limited to a particular protein, and the level of recognition of different proteins varied among the six patients.

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References De Maria et al. 1997

- CD3+ cells that also carry a natural killer cell receptor (NKR+) can exhibit down regulation of T cell function.
- · Anti-NKR IgM MAb masked this inhibitory function and increased HIV-1 specific CTL activity in phytohemagglutininactivated PBMC cultured in the presence of IL-2 from 3/5 patients, and in one other case anti-NKR MAb brought HIV-1 specific CTL activity to detectable levels.

HXB2 Location Pol

Author Location Pol (LAI, MN)

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

References Goh et al. 1999

- 13/37 exposed uninfected individuals with repeated high-risk sexual exposure had HIV-1 specific CTL against Env, Gag, Pol, or a combination of proteins - CTL activity was correlated with a CCR5 wildtype genotype.
- In this group, the highest CTLp frequencies were directed at Gag, but the most common response was to Env and four individuals had responses to multiple HIV-1 proteins.

HXB2 Location Pol

Author Location Pol (LAI)

**Epitope** Subtype B Immunogen vaccine

Vector/Type: canarypox HIV component:

Gag, gp120, gp41, Nef, Protease, RT

Species (MHC) human

References Evans et al. 1999

• A Canarypox vaccine expressing gp120, gp41, Gag, Protease, Nef and Pol CTL epitopes gave rise to CTL that could be detected in 61% of the volunteers - responses to Gag, Env, Nef and Pol were detected 3-6 months after the last vaccination.

HXB2 Location Pol

Author Location Gag/Pol (MN)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env,

Gag, Pol Adjuvant: CD80, CD86

Species (MHC) chimpanzee

References Kim et al. 1998

· The study explores the use of co-stimulatory molecules coexpressed with an HIV-1 immunogen in a DNA vaccine to enhance the immune response - co-expression of CD86, but not CD80, dramatically increased both HIV Env and Gag/Pol specific CTL and Th proliferative responses.

HXB2 Location Pol

**Author Location** Pol (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Jin et al. 1998a

• CTL precursor frequencies were determined in HIV-1 infected pregnant women, and significantly higher CTLp frequencies to Pol and Nef were found in non-transmitting mothers than in transmitting mothers;

HXB2 Location Pol

**Author Location** Pol

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Young et al. 2001

- Addition of recombinant rec human IL12 (rhIL12) to cultures increased HIV-specific lysis of HIV-Gag, Pol and gp120 vaccinia expressed antigens (11/15 tested increased lysis by > 5%) if the culture was derived from HIV+ individuals who had CD4 cells/ul > 500.
- 2/10 individuals with <200 CD4 cells/ul, and 3/10 individuals with 200-500 CD4cells/ul, had an increase of >5% upon treatment of the culture with rhIL12, so a few individuals in late stage disease had CD8 cells that maintained responsiveness to rhIL12.

HXB2 Location Pol

Author Location RT (subtype A, B, D)

**Epitope** 

Subtype A, B, D

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons References Cao et al. 2000

- HIV-1 subtypes A and D dominate the Ugandan epidemic, and a vaccine trial using B clade antigen is underway – this study addresses relative levels of cross-reactive CTL responses in HIV infected Ugandans to A, D, and B clade recombinant vaccinia viruses expressing Gag, Env, Pol, RT or Nef from HIV-1 clades A, B, and D.
- Proteins corresponding to the subtype of the infecting strains tended to trigger higher levels of CTL response measured by percent specific lysis, but there was extensive inter-subtype cross-reactivity with B clade proteins and the co-circulating subtype.

HXB2 Location Pol Author Location Pol

Epitope

Immunogen HIV-1 infection

Species (MHC) human

References White et al. 2001

 HIV-specific CTL activity was detected in the female reproductive tract of only 1/3 HIV-infected women who underwent a hysterectomy, although CTL could be identified in the PBMC of all three women.

HXB2 Location Pol

Author Location Pol (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Jin *et al.* 2000a

- The CTL precursor level (CTLp) was measured in long term non-progressors (LTNP) with low viral load using limiting dilution analysis and measuring CTL against Env Gag and Pol expressed in vaccinia in autologous targets.
- LTNPs have high memory CTL numbers and low viral load.

HXB2 Location Pol

Author Location Pol

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

**Keywords** review, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 2001

- This is a review that summarizes observations about HIV-specific CTL found in the HIV-1 exposed persistently seronegative (HEPS) population.
- The CTL responses assayed by ELISPOT and by CTL precursor frequencies by limiting dilution analysis indicate that CTL in HEPS individuals tend to be of a lower magnitude than in chronic HIV-1 infections the responses in HEPS cases are below the level of detection by tetramer assays.
- CD8+ CTL responses tend to be detectable in HEPS subjects only if they are recently exposed, and the response diminishes if exposure is reduced it is not clear if there is a stable memory population in HEPS cases.

- CD8+ CTL responses in the HEPS population are associated with HIV-1 specific CD4+ T cell responses, assayed by proliferation assays, IL-2 secretion, and ELISPOT, and the authors consider the possibility that HIV-1-specific T-help responses improve the "quality" of the CD8+ response in HEPS individuals relative to HIV-1 infected individuals, who tend to have a poor HIV-1-specific T-help response.
- HIV-1 specific CD8+ CTL responses in HIV-1 infected individuals show reduced levels of perforin, and the T cells may not mature properly, and although similar studies have not been conducted in HEPS individuals this is considered as a possible difference in the CTL immune response in HEPS and HIV-1 infected people.

HXB2 Location Pol

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS), responses in children, mother-to-infant transmission

References De Maria et al. 1994; Kuhn et al. 2002

- 6/24 HIV uninfected infants (ages 15-50 months) born to HIV+ mothers had HIV-1 specific CTL responses to vacciniaexpressed Nef, Gag/Pol, Env.
- Reviewed in Kuhn et al. [2002].

**HXB2 Location** Pol

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, HIV exposed persistently seronegative (HEPS), responses in children, mother-to-infant transmission, rate of progression

References Kuhn et al. 2002; Wasik et al. 1999

- In HIV-infected infants HIV-specific, CTL responses were not detectable in icord blood or in PBMC collected shortly after birth and were absent or remained very low in 3 infants with a rapidly progressive disease. For those who progressed more slowly, the HIV-specific CTL activity varied.
- The two infants with high levels of Env peptide-stimulated IL-2 responses had the highest CTLp frequencies.
- Stronger responses were detected after initiation of the antiretroviral therapy.
- Two babies that were not infected though born to HIV+ mothers had detectable though low HIV-specific CTLp responses to Env (1/2), Pol (2/2), Gag (1/2) cord blood and transiently in PBMC after birth.
- Reviewed in Kuhn et al. [2002].

HXB2 Location Pol Author Location

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS), responses in children, mother-to-infant transmission

References Aldhous et al. 1994; Kuhn et al. 2002

- Six of nine HIV vertically infected infants had HIV-1 specific CTL responses to vaccinia expressed Tat (4/6), Pol (6/6), Env (1/6), or Gag (1/6), but not all responses were detected at all time points.
- Two of eleven babies that were not infected though born to HIV+ mothers had detectable responses Tat (1/2), Pol (2/2), Gag (1/2).
- Reviewed in Kuhn et al. [2002].

HXB2 Location Pol

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed; however, epitopes were not found that span the invariant, most highly conserved regions of RT and Protease. This might be due to the virus evolving conserved features that disallow the CTL responses in these most conserved regions, as functional constraints for enzyme function would not tolerate change, and normal capacity for immune escape by rapid evolution is lost in these domains.

HXB2 Location Pol

**Author Location** 

**Epitope** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, subtype comparisons

References Loemba et al. 2002

• Therapeutic RT inhibitors were used to select *in vitro* for resistance mutations in subtype C viruses. Many of the resistance mutations were located within analogs to CTL epitopes that had been defined for the B subtype,

HXB2 Location Pol

**Author Location (IIIB)** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, acute/early infection

References Ortiz et al. 2002

 Subjects treated with HAART early in HIV-infection showed a correlation between the number of viremic episodes and the total as well as the Pol-specific CD8 T-cell activity as measured by Elispot SFC per million PBMC summed across Pol, Env, Nef and Gag. The subjects treated early after infection had higher levels of CD8+ T-cell activity (N = 31) than those treated later (N = 23), and a greater capacity to enhance CD8+ T-cell responses to viremic episodes.

HXB2 Location Pol

**Author Location** (MN)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression

References Edwards et al. 2002

- 96% (26/27) chronically infected HIV-1 infected patients elicited gamma-IFN CD8+ T-cell responses against Gag.
- Nef and/or Pol CTL responses were detected in 86% of the subjects.
- The magnitude and breadth of Gag and p24 T-cell responses correlated with absolute CD4 counts, and inversely correlated with viral load.
- Pol and Int CTL responses correlated positively with absolute CD4+ T-cell count.
- Nef and Env responses did not correlate with either CD4 counts or viral load.

HXB2 Location Pol

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, dendritic cells

References Larsson et al. 2002b

 Autologous mature dendritic cells with rec vaccinia expressing Gag, Pol, Nef and Env could amplify CD8+ T-cell Elispot responses 4-38 fold in five HIV+ patients on successful HAART treatment, relative to autologous monocytes. Some weak responses could only be detected using mature dendritic cells as APCs, and this approach could be useful for detection of low frequency memory cells.

HXB2 Location Pol

**Author Location** (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords immunotherapy

References Trickett et al. 2002

 Conditions were optimized for ex-vivo expansion of CD8+ and CD4+ T-cells with the goal of functional T-cell production for autologous immunotherapy. 10,000-fold expansions were obtained in 14 days with optimized concentrations of IL-2, anti-CD3 and anti-CD28 coated microspheres, and decreasing amounts of serum over the first 8 days.

HXB2 Location Pol

**Author Location** (IIIB)

**Epitope** 

Subtype B

Immunogen HIV-1 and HCV co-infection

Species (MHC) human

**Keywords** rate of progression

References Lauer et al. 2002

- HIV-1 and HCV immune responses were studied in 22 individuals who were co-infected with HIV-1 and hepatitis C virus (HCV). IFNgamma production was measured in an Elispot assay of CD8+ T-cells using targets expressing either Gag, RT, Env and Nef in a vaccinia construct, or one of seven HCV proteins.
- All 22 patients targeted at least one protein. 20/22 patients recognized RT, 17/22 patients recognized Gag, 13/22 subjects recognized Env and 11/22 patients recognized Nef. Robust CTL activity was independent of disease progression or viral load
- Despite high HCV viral loads, very few HCV CD8+ T-cell Elispot responses were detected. In a control HCV infected person who did not have HIV-1, strong anti-HCV responses were mounted.
- HIV-specific CD4 proliferative responses were detected in 9/17 coinfected patients, but no HCV responses were detected.

HXB2 Location Pol Author Location Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, responses in children **References** Scott *et al.* 2001

- CTL responses before and after initiation of ART were studied in 13 HIV-1 vertically infected infants <6 months of age, and 4 that were >6 months of age.
- Before ART 2/13 infants <6 months of age showed IFNgamma Elispot CD8+ T-cell responses, one to Nef and one to Env and Nef, and these responses became undetectable after successful therapy—3 infants were coinfected with CMV and all 3 had CMV-specific CD8+ T-cell responses.
- One older infant, at 23 months, had CTL responses against all for proteins tested, Gag, Pol, Nef and Env, and had the lowest plasma viremia of the study group. 3/4 infants older than 6 months of age responded to either Nef or Pol.
- Administration of ART over 48 weeks broadened the HIV-1specific CTL response in 2/4 of the older children that were incomplete responders.

HXB2 Location Pol Author Location (IIIB, MN)

> Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords dendritic cells

References Larsson et al. 2002a

 Dendritic cells acquire and present HIV-1 antigens derived from dead, apoptotic cells or from non-infectious, fusioncompetent HIV-1 virions, and these DC cells could stimulate CD4+ and CD8+ T-cells resulting in IFNgamma production in an Elispot assay. Both HLA Class I and class II molecules were used for presentation. This may be an important aspect of the initial immune response to HIV-1 infection of CD4+ cells in the mucosal subepithelia.

HXB2 Location Pol

**Author Location** (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Ortiz et al. 2001

 Immune responses in eight chronically HIV-1 infected patients undergoing HAART therapy structured treatment interruptions (STI) were studied. STI boosted HIV-1 specific CTL responses and elevated CTL responses were maintained up to 22 weeks after the last treatment interruption, but viral load rebound to pretreatment levels and CD4 T-cell count decline was observed. CD8 responses in PBMC were measured by cytokine flow cytometry with gp160, Gag p55, RT-Pol and Nef expressed in vaccinia.

## II-B-16 Vif CTL/CD8 + epitopes

**HXB2 Location** Vif (17–26)

**Author Location** (LAI)

**Epitope** RIRTWKSLVK

Subtype B

Immunogen

Species (MHC) human (A\*0301)

Keywords optimal epitope

References Altfeld 2000; Frahm et al. 2007

**HXB2 Location** Vif (17–26)

**Author Location** Vif (17–26 SF2)

**Epitope** RIRTWKSLVK

Epitope name RK10

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

References Altfeld et al. 2001a

- CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- 10/29 (35%) individuals tested responded to Vif.
- This epitope was recognized by 3/15 individuals expressing A\*0301 allele.
- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- Overlapping Vif peptides QVDRMRIRTWKSLVK and RIRTWKSLVKHHMYI both reacted with T-cells from AC-06 and contained epitope RIRTWKSLVK.

**HXB2 Location** Vif (17–26)

**Author Location** Vif (17–26)

Epitope RIRTWKSLVK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vif (17-26)

**Author Location** 

**Epitope** RIRTWKSLVK Epitope name Vif-RK10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3) References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA A03, 3/21 (14%) recognized this epitope.

HXB2 Location Vif (17-26) **Author Location** Vif (17–26)

Epitope RIRTWKSLVK

**Epitope name** A3-RK10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 1/7 individuals had detectable responses to this epitope after

HXB2 Location Vif (17–26)

**Author Location** Vif (17–26)

Epitope RIRTWKSLVK

Epitope name A3-RK10 Vif

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords supervised treatment interruptions (STI), es-

cape, early treatment, superinfection

References Altfeld et al. 2002a

- An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.
  - The second infecting strain had the variant riStwkslvk. The initial CTL response to persisted to against both variants after the superinfection was established.

HXB2 Location Vif (17-26)

**Author Location** Vif (17–26)

Epitope RIRTWKSLVK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

HXB2 Location Vif (17–26)

**Author Location** (B consensus)

Epitope RIRTWKSLVK

Epitope name RK10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A02, A03, B08, B62, Cw7, Cw10; A03, B07, Cw7

Assay type Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.
- 2/9 individuals recognized this epitope, presented by HLA-A3.

**HXB2 Location** Vif (17–26)

**Author Location** Vif

**Epitope** RIRTWKSLVK

**Epitope name** RK10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 6, rirtwNslvk, was found not to correspond to the most polymorphic residue in the epitope.

**HXB2 Location** Vif (17–26)

**Author Location** 

Epitope RIRTWKSLVK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Vif (23-31)

**Author Location** Vif (23–)

**Epitope** SLVKHHMYV **Epitope name** Vif23(9V)

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Vif Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (MHC) human, transgenic mouse (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release

assay, Flow cytometric T-cell cytokine assay **Keywords** binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced CTL and CD8+ T-cell IFN gamma responses in mice. Response was detected in 1/17 HIV+ HLA-A2 subjects.
- The variant slvkhhmyI was an intermediate A2 binder, and stimulated immune reponses in fewer A2 transgenic mice. The same person recognized both variants.

HXB2 Location Vif (27–41)

**Author Location** Vif

Epitope HHMYISKKAKGWGWFYR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot 33% (23/70) targeted one or more Vif peptides, and this peptide was the most frequently recognized epitope in Vif (25%).
- The regulatory proteins Rev and Tat combined contributed to 3%, and the accessory proteins Vif, Vpr and Vpu to 7%, of the total magnitude of HIV-1 specific CTL responses in a subset of 22 HIV-1 infected individuals in whom all HIV-1 proteins were studied.

HXB2 Location Vif (28-36)

**Author Location** Vif (28–36)

Epitope HMYISKKAK

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (A\*0301)

**Keywords** early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vif (28–36)

**Author Location** Vif (28–36)

Epitope HMYISKKAK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

HXB2 Location Vif (28–36) Author Location Vif (28–36) Epitope HMYISKKAK Epitope name A3-HK9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3) Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection.
   2/7 individuals had detectable responses to this epitope after STI.

HXB2 Location Vif (31–39)
Author Location Vif (31–39 SF2)
Epitope ISKKAKGWF
Immunogen HIV-1 infection
Species (MHC) human (B\*5701)
References Altfeld *et al.* 2001a

- CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- 10/29 (35%) individuals tested responded to Vif.
- $\bullet$  This epitope was recognized by 2/6 individuals expressing  $B \! * \! 5701$  allele.

HXB2 Location Vif (31–39) Author Location Vif (31–39) Epitope ISKKAKGWF

Subtype B

Immunogen HIV-1 infection
Species (MHC) human (B\*5701)
Keywords early-expressed proteins
References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

**HXB2 Location** Vif (31–39) **Author Location** Vif (31–39)

Epitope ISKKAKGWF

Immunogen

Species (MHC) human (B\*5701) Keywords optimal epitope References Frahm *et al.* 2007

HXB2 Location Vif (31–39)

**Author Location** Vif

Epitope VSKKAKGWI

Epitope name VI9 Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 1,Askkakwi, was found in the most polymorphic residue in the epitope.

HXB2 Location Vif (31-39)

**Author Location** 

Epitope ISKKAKGWF
Immunogen HIV-1 infection
Species (MHC) human (B57, B\*5801)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** responses in children, mother-to-infant transmission, escape

References Feeney et al. 2005

• Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.

HXB2 Location Vif (31–39)

**Author Location** 

Epitope ISKKAKGWF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** epitope processing, escape **References** Yusim *et al.* 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

**HXB2 Location** Vif (41–57) **Author Location** (C consensus)

Epitope RHHYESRHPKVSSEVHI

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*1510)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Vif (48–57) **Author Location** Vif (48–57 SF2)

**Epitope** HPRVSSEVHI

Epitope name HI10

Immunogen HIV-1 infection Species (MHC) human (B\*0702) References Altfeld *et al.* 2001a

- CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- 10/29 (35%) individuals tested responded to Vif.
- This epitope was recognized by 3/8 individuals expressing B\*0702 allele.
- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- Overlapping Vif peptides HHYESTHPRVSSEVH and TH-PRVSSEVHIPLG both reacted with T-cells from AC-06 and contained epitope HPRVSSEVHI.

HXB2 Location Vif (48–57)

**Author Location** Vif (48–57)

**Epitope** HPRVSSEVHI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

**Keywords** early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vif (48-57)

**Author Location** Vif (48–57)

Epitope HPRVSSEVHI

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Vif (48–57)

**Author Location** (C consensus)

Epitope HPKVSSEVHI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** Vif (48–57)

Author Location (C consensus)

Epitope HPKVSSEVHI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the H1 residue of HPKVSSEVHI are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** Vif (48–57)

**Author Location** Vif (48–57)

Epitope HPRVSSEVHI

Epitope name B7-HI10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7) Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 2/4 individuals had detectable responses to this epitope after STI.

**HXB2 Location** Vif (48–57)

**Author Location** Vif (48–57)

Epitope HPRISSEVHI

Epitope name B7-HM0 Vif

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFNγ

Keywords supervised treatment interruptions (STI), es-

cape, early treatment, superinfection

References Altfeld et al. 2002a

- An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.
- The second infecting strain had the variant hpKissevhi. The CTL response was equal against both variants, and declined over time.

HXB2 Location Vif (48–57)

**Author Location** (B consensus)

Epitope HPRISSEVHI

Epitope name HI10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A03, B07, Cw7

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

HXB2 Location Vif (48-57)

**Author Location** (B consensus)

**Epitope** HPKISSEVHI

**Epitope name** HKI10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A03, B07, Cw7

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

HXB2 Location Vif (48-57)

**Author Location** Vif

**Epitope** HPRISSEVHI

Epitope name HI10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 2, hSrissevhi, was found not to correspond to the most polymorphic residue in the epitope.

**HXB2 Location** Vif (48–57)

**Author Location** 

Epitope HPRVSSEVHI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

**HXB2 Location** Vif (54–63)

Author Location (C consensus)

Epitope EVHIPLGEAR

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the R10 residue of EVHIPLGEAR are associated with the presence of the HLA presenting molecule in the host.
- EVHIPLGEAR not optimized.

HXB2 Location Vif (57–66)

**Author Location** Vif (57–66)

Epitope IPLGDAKLII

Immunogen

Species (MHC) human (B51)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Vif (57–66)

**Author Location** Vif (57–66)

Epitope IPLGDAKLII

Epitope name II10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B51)

Donor MHC A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, Chromium-release assay

Keywords escape, acute/early infection

References Bansal et al. 2005

- Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- The peptide that carries this epitope was recognized at high levels early in infection, and the response to this epitope diminished over time. The epitope sequence varied between months 3 and 32.

**HXB2 Location** Vif (61–69)

**Author Location** Vif (61–69)

Epitope DAKLIITTY

Epitope name DY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

**Donor MHC** A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, Chromium-release assay

**Keywords** escape, acute/early infection

References Bansal et al. 2005

- Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- The peptide that carries this epitope was recognized early in infection but the response diminished over time. A point mutation of epitope position 7 (T to K, DAKLIIkTY) was detected at high frequency at a chronic infection timepoint, 34 months.

The K variant was an escape form, and had low avidity by gamma IFN Elispot.

**HXB2 Location** Vif (61–80) **Author Location** Vif (61–80)

Epitope EARLVIKTYWGLOTGERDWH

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** Vif (71–90) **Author Location** Vif (71–90)

Epitope GLQTGERDWHLGHGVSIEWR

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Vif (72–89) Author Location (C consensus)

Epitope LQTGERDWHLGHGVSIEW

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*5703)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Vif (73–81) **Author Location** Vif (73–81)

Epitope HTGERDWHL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*35)

**Donor MHC** A\*03, A\*24, B\*35, B\*40

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** acute/early infection, variant crossrecognition or cross-neutralization, superinfection

References Yang et al. 2005b

- An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- The response to this epitope, HTGERDWHL, was present before superinfection but waned afterward. The epitope from the first strain had the substitution hPgerdwhl, while the second strain matched the test peptide.

HXB2 Location Vif (79–87)
Author Location Vif (79–87)
Epitope WHLGHVSI
Immunogen HIV-1 infection
Species (MHC) human (B\*1510)

Keywords optimal epitope

**References** Frahm *et al.* 2007

HXB2 Location Vif (79–87) Author Location Nef (C consensus) Epitope WHLGHGVSI

Epitope name WI9 Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*1510)

**Donor MHC** A\*2601, A\*7401, B\*0801, B\*1510, Cw\*0202, Cw\*0801

Cw 0202, Cw

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords assay standardization/improvement, charac-

terizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was one of two used to illustrate how specific epitopes were characterized with regard to defining the optimal epitope and the HLA restricting element. HLA allelic associations in the population with peptide recognition was generally high predictive of the epitope within the 15 mer.

HXB2 Location Vif (79–87)

**Author Location** (C consensus)

Epitope WHLGHGVSI

Subtype C

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*1510)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location Vif (79–87)

**Author Location** (C consensus)

**Epitope** WHLGHGVSI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1510)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords epitope processing, rate of progression, opti-

mal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the V7 residue of WHLGHGVSI are associated with the presence of the HLA presenting molecule in the host. Mutation of a residue outside of the optimized epitope is also associated with the HLA.

**HXB2 Location** Vif (79–87)

**Author Location** Vif (79–87)

Epitope WHLGHGVSI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1510)

Assay type Other

Keywords epitope processing, HLA associated polymor-

phism

References Boutwell & Essex 2007

• All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.

 WHLGHGVSI was a previously defined B\*1510 presented epitope that was associated with a polymorphism, dlWHLGHGVSI,in the first position before that epitope. This epitope was embedded in a previously identified CTL immunoreactive region.

HXB2 Location Vif (79–87)

**Author Location** Vif (79–87)

Epitope WHLGQGVSI

Immunogen HIV-1 infection

Species (MHC) human (B\*3801)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Vif (85–102)

Author Location (C consensus)

Epitope VSIEWRLRRYSTQVDPGL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*18)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Vif (101–109)

**Author Location** Vif (101–)

Epitope GLADQLIHL

**Epitope name** Vif101(9L)

**Immunogen** HIV-1 infection, vaccine, computer prediction *Vector/Type:* peptide *Adjuvant:* Incomplete

Freund's Adjuvant (IFA)

Species (MHC) human, transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder that induced CTL and CD8+ T-cell IFN gamma responses in mice. Responses were detected in 3/17 HIV+ HLA-A2 subjects.
- The variant gladqlihM was an intermediate A2 binder, but still could stimulate a response in HLA-A2 transgenic mice.
   It was not recognized by the 3 people who recognized with GLADQLIHL.

HXB2 Location Vif (101–110)

**Author Location** Vif

Epitope DLADQLIHLY

Subtype multiple
Immunogen HIV-1 infection
Species (MHC) human (A2)

**Donor MHC** A02, A30, B39; A01, A02, B08, Cw16

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

References De Groot et al. 2003

Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.

• Estimated binding probability for DLADQLIHLY: 54%

HXB2 Location Vif (102–111)

Author Location Vif (102–111 SF2)

Epitope LADQLIHLHY

Immunogen HIV-1 infection

Species (MHC) human (B\*1801)

References Altfeld *et al.* 2001a

- CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- 10/29 (35%) individuals tested responded to Vif.
- This epitope was recognized by 2/5 individuals expressing B\*1801 allele.

Author Location Vif (102–111)
Epitope LADQLIHLHY
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (B\*1801)
Keywords early-expressed proteins

**HXB2 Location** Vif (102–111)

**References** Addo *et al.* 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vif (102–111)
Author Location Vif (102–111)
Epitope LADQLIHLHY
Immunogen HIV-1 infection
Species (MHC) human (B\*1801)
Keywords optimal epitope
References Frahm et al. 2007

HXB2 Location Vif (102–111)
Author Location Vif (102–110)
Epitope LADQLIHLHY
Epitope name LY10
Subtype B

Immunogen HIV-1 infection Species (MHC) human (B18)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, Flow cytometric T-cell cytokine

**Keywords** optimal epitope **References** Allen *et al.* 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- This epitope did not vary.

HXB2 Location Vif (102–111)
Author Location Vif (102–110)
Epitope LADQLIHLHY
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (B18)
Donor MHC A2, A11, B18, B44, Cw5, Cw12
Country United States.
Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** Vif (102–111) **Author Location** 

Epitope LADQLIHLHY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated, traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.

- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
   CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

**HXB2 Location** Vif (106–123) **Author Location** (C consensus)

Epitope LIHMHYFDCFADSAIRKA

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*6801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Vif (127–135)

**Author Location** Vif

**Epitope HIVSPRCEY** 

Epitope name HY9 Subtype B

Immunogen HIV-1 infection Species (MHC) human (A29)

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 3, HIgSPRCEY, was found not to correspond to the most polymorphic residue in the epitope.

**HXB2 Location** Vif (127–135)

**Author Location** Vif (125–135)

Epitope HIVSPRCEY

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** 1261: A\*0201, A29, B58, B62, Cw\*0304,

Cw\*1601

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

**HXB2 Location** Vif (149–157)

**Author Location** Vif (149–)

**Epitope** ALAALITPK

**Epitope name** Vif149

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Vif Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (MHC) human, transgenic mouse (A2)

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced CTL and CD8+ T-cell IFN gamma responses in mice. Responses were detected in 1/17 HIV+ HLA-A2 subjects.

**HXB2 Location** Vif (151–168)

**Author Location** (C consensus)

Epitope TALIKPKKIKPPLPSVRK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1701)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Vif (158–166)

Author Location Vif (158–) Epitope KIKPPLPSV Epitope name Vif158(2I)

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Vif Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human, transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced CD8+ T-cell IFN gamma responses in mice. Responses were detected in 1/17 HIV+ HLA-A2 subjects.
- The substitution kTkpplpsv was also a good binder, but did not elicit a response in transgenic mice, and no reponse to this variant was detected among the 17 HIV+ people tested.

**HXB2 Location** Vif (158–168)

**Author Location** Vif (158–168)

Epitope KTKPPLPSVKK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

**HXB2 Location** Vif (158–168)

**Author Location** Vif (158–168)

Epitope KTKPPLPSVKK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

**Keywords** optimal epitope

References Frahm et al. 2007

**HXB2 Location** Vif (158–168)

Author Location (B consensus)

**Epitope** KTKPPLPSVKK

Epitope name KK11

Immunogen HIV-1 infection Species (MHC) human (A11)

**Donor MHC** A02, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** Vif (158–168)

**Author Location** Vif (158–168)

**Epitope** KTKPPLPSVKK

Epitope name A3-KK11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 2/7 individuals had detectable responses to this epitope after STI.

**HXB2 Location** Vif (158–168)

**Author Location** Vif (158–168)

**Epitope** RRKPPLPSIAK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** supervised treatment interruptions (STI), es-

cape, superinfection

References Altfeld et al. 2002a

An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response to 25 distinct epitopes, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.

• The second infecting strain had the variant rTkpplpsVTk. The patient maintained persistent reactive CTL against both variants after the superinfection was established.

**HXB2 Location** Vif (158–168)

Author Location Vif

**Epitope** RIKPPLPSVTK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- RIKPPLPSVTK had to mutations overtime, in positions 1 and 10: KikpplpsvKk

**HXB2 Location** Vif (158–168)

**Author Location** Vif

**Epitope** KIKPPLPSVTK

Epitope name KK11 Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 10, KIKPPLPSVkK, was found in the most polymorphic residue in the epitope.

**HXB2 Location** Vif (160–169)

**Author Location** Vif

Epitope KPPLPSVKKL

Immunogen

Species (MHC) human (B7)

References De Groot et al. 2001

- The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.
- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes could stimulate IFNγ production in an ELISPOT assay.

 KPPLPSVKKL was newly identified as an HLA-B7 epitope in this study.

**HXB2 Location** Vif (160–169)

**Author Location** Vif

Epitope KPPLPSVKKL

Epitope name 1296

Subtype multiple

Immunogen HIV-1 infection Species (MHC) human (B7)

Donor MHC A03, A24, B07, B38, Cw07, Cw12/13

Country United States.

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope prediction

**References** De Groot *et al.* 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for KPPLPSVKKL: 23%

**HXB2 Location** Vif (168–176)

**Author Location** Vif

Epitope KLTEDRWNK

Epitope name 1344

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A03, A24, B27, B57, Cw13, Cw18

Country United States.

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope prediction

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for KLTEDRWNK: 54%

**HXB2 Location** Vif

**Author Location** Vif

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Nef, Vif, Vpu

Species (MHC) mouse (H-2<sup>d</sup>)

Keywords subtype comparisons, Th1

References Ayyavoo et al. 2000

 Splenocytes from BALB/c mice immunized with pVVN-P DNA were incubated with Vif, Vpu or Nef antigens for 3 days and assayed for IL-4 and IFN-gamma levels.

- Antigen stimulation increased IFN-gamma production in pVVN-P immunized mice, indicating a Th1 response.
- IL-4 production was not significantly changed after antigen stimulation compared to control levels.
- Cross-clade CTL activity was also observed: A, B clade, CRF01(AE) clade antigens could serve as targets for the B clade immunization-stimulated CTL - an HIV-1 AC recombinant, however, did not stimulate a CTL response, but was expressed at lower levels on the target cell.

**HXB2 Location** Vif Author Location Vif **Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Nef, Author Location Tat Vif, Vpu

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords subtype comparisons, Th1

References Ayyavoo et al. 2000

- Splenocytes from BALB/c mice immunized with pVVN-P DNA were incubated with Vif, Vpu or Nef antigens for 3 days and assayed for IL-4 and IFN-gamma levels.
- · Antigen stimulation increased IFN-gamma production in pVVN-P immunized mice, indicating a Th1 response.
- IL-4 production was not significantly changed after antigen stimulation compared to control levels.
- Cross-clade CTL activity was also observed: A, B clade, CRF01(AE) clade antigens could serve as targets for the B clade immunization-stimulated CTL - an HIV-1 AC recombinant, however, did not stimulate a CTL response, but was expressed at lower levels on the target cell.

**HXB2 Location** Vif **Author Location** Vif

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env, Gag, Pol, Vif Adjuvant: B7, IL-12

Species (MHC) mouse

References Kim et al. 1997c

- · A gag/pol, vif or env DNA vaccine, when delivered in conjunction with the plasmid encoding the co-stimulatory molecules B7 and IL-12, gave a dramatic increase in both the cytotoxic and proliferative responses in mice.
- When IL-12 was present, CTL response could be detected even without in vitro stimulation.

## II-B-17 Vpr CTL/CD8 + epitopes

HXB2 Location Vpr (1–18)

**Author Location** Vpr

Epitope MEQAPENQGLQREPYNEW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Novel unmapped epitope. A mutation occurred over time in an individual that reacted to this peptide: MEQAPENQGpQREPYNEW.

HXB2 Location Vpr (9-20)

Epitope APQGHPNNQVSI

**Epitope name** AI12 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A2, B7, B44, Cw5, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- · Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 4, APQdHPNNQVSI, was found not to correspond to the most polymorphic residue in the epitope. This is a novel unmapped epitope.

HXB2 Location Vpr (9-26)

Author Location (C consensus)

Epitope GPQREPYNEWTLELLEEL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0704)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

Keywords rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location Vpr (12-20)

**Author Location** Vpr (12–20 SF2)

Epitope REPHNEWTL

Immunogen HIV-1 infection

Species (MHC) human (B\*4002)

Keywords acute/early infection

References Altfeld et al. 2001a

- CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- Individuals with long-term nonprogressive and treated chronic HIV-1 infection targeted Vpr more frequently than individuals with treated acute infection.
- Only one B\*4002+ individual was tested, and had a CTL response against REPHNEWTL.
- Vpr is a frequent target of HIV-1 specific CD8+ T-cells a response was detected in 45% of individuals tested and Vpr and p17 were the most preferentially targeted proteins per unit length by CD8+ T-cells.

**HXB2 Location** Vpr (12–20) **Author Location** Vpr (12–20)

Author Location vpr (12–20)

 ${\bf Epitope} \ \ {\tt REPHNEWTL}$ 

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*4002)

**Keywords** early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vpr (12–20)

**Author Location** 

Epitope REPHNEWTL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated, traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Vpr (19-28)

**Author Location** Vpr

**Epitope TLEILEELKN** 

Epitope name TN10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3?)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One mutation, at position 1, Aleileelkn,occurred over time. This is a novel unmapped epitope.

HXB2 Location Vpr (25-40)

Author Location Vpr (25-40 HXB2)

Epitope ELKNEAVRHFPRIWLH

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- Responses to this peptide were detected in 17% of the study subjects, and it was one of the 25 most frequently recognized peptides.

HXB2 Location Vpr (29–37)

**Author Location** Vpr (29–37 2001 HIV-1 subtype B cons)

Epitope EAVRHFPRI

Epitope name EI9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

Donor MHC A\*0201. A\*2501. B\*1801. B\*5101. Author Location Vpr

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- Last position (9)in the epitope had potentially experienced positive selection. EAVRHFPRt and EAVRHFPRl escape variants were found.

HXB2 Location Vpr (29–37)

Author Location Vpr (29-37)

Epitope EAVRHFPRI

Immunogen

Species (MHC) human (B51)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Vpr (29–37)

**Author Location** Vpr (29–37 B)

**Epitope** EAVRHFPRI

Immunogen HIV-1 infection

Species (MHC) human (B51)

**Donor MHC** A\*0201, A\*2501, B18, B51, Cw\*0102,

Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFNsecreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- · All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

HXB2 Location Vpr (29–37)

**Epitope** EAVRHFPRI

Epitope name EL9

Immunogen

Species (MHC) (B51)

Keywords review, immunodominance, acute/early infection, early-expressed

proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

• This review discusses the importance of 3 factors that impact the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides listed as immunodominant in acute HIV-1 infection.

HXB2 Location Vpr (30-38)

**Author Location** Vpr (29–38 SF2)

Epitope AVRHFPRIW

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords acute/early infection

References Altfeld et al. 2001a

- CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- This epitope was recognized by 4/6 individuals expressing B\*5701 allele.
- Individuals with long-term nonprogressive and treated chronic HIV-1 infection targeted Vpr more frequently than individuals with treated acute infection.
- Vpr is a frequent target of HIV-1 specific CD8+ T-cells a response was detected in 45% of individuals tested and Vpr and p17 were the most preferentially targeted proteins per unit length by CD8+ T-cells.

HXB2 Location Vpr (30–38)

Author Location Vpr (29–38)

Epitope AVRHFPRIW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

**Keywords** early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vpr (30–38)

Author Location Vpr (30–38)

Epitope AVRHFPRIW

Immunogen

Species (MHC) human (B\*5701)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Vpr (30–38)

**Author Location** 

Epitope AVRHFPRIW Epitope name Vpr-AW9 Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)
References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA B57, 1/7 (14%) recognized this epitope.

**HXB2 Location** Vpr (30–38) **Author Location** Vpr (30–38)

Epitope AVRHFPRIW

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B57)

Donor MHC A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

HXB2 Location Vpr (30–38)

**Author Location** 

Epitope AVRHFPRIW
Immunogen HIV-1 infection
Species (MHC) human (B57, B\*5801)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

Keywords responses in children, mother-to-infant trans-

mission, escape

References Feeney et al. 2005

• Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.

HXB2 Location Vpr (30-38)

**Author Location** Vpr

Epitope AVRHFPRIW

Subtype B, C

Immunogen HIV-1 infection Species (MHC) human (B57, B63)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This epitope was recognized by 30% of B63-positive subjects and 14% of B57/58-positive subjects.

HXB2 Location Vpr (30–38)

**Author Location** 

Epitope AVRHFRPIW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated, traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Vpr (31–39)

Author Location Vpr (31-39)

Epitope VRHFPRIWL

Immunogen HIV-1 infection

Species (MHC) human (B27)

**Keywords** optimal epitope

References Frahm et al. 2007

HXB2 Location Vpr (31–50)

**Author Location** Vpr (31–50)

Epitope VRHFPRPWLHSLGQYIYETY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons

References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 8 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Vpr (34–42)

Author Location Vpr (34–)

Epitope FPRPWLHGL

Epitope name Vpr34

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Vpr Adjuvant: Incomplete Freund's Adjuvant (IFA)

**Species (MHC)** human, transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced a CD8+ T-cell IFN gamma response in 1/6 mice. Responses were detected in 3/17 HIV+ HLA-A2 subjects.

HXB2 Location Vpr (34-42)

**Author Location** Vpr (34–42 SF2)

Epitope FPRIWLHGL

Epitope name FL9

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Keywords acute/early infection

References Altfeld et al. 2001a

- CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- This epitope was recognized by 2/2 individuals expressing B\*8101 allele and 4/8 individuals expressing B\*0702 allele.
- Individuals with long-term nonprogressive and treated chronic HIV-1 infection targeted Vpr more frequently than individuals with treated acute infection.
- Vpr is a frequent target of HIV-1 specific CD8+ T-cells a response was detected in 45% of individuals tested and Vpr and p17 were the most preferentially targeted proteins per unit length by CD8+ T-cells.
- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.

• FPRIWLHGL was the only epitope identified in Vpr for AC-

HXB2 Location Vpr (34–42)

Author Location Vpr (34-42)

Epitope FPRIWLHGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vpr (34–42)

**Author Location** Vpr (34–42)

Epitope FPRIWLHGL

**Immunogen** HIV-1 infection

Species (MHC) human (B\*0702)

**Keywords** optimal epitope

References Frahm et al. 2007

HXB2 Location Vpr (34-42)

**Author Location** Vpr

Epitope FPRIWLHGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Donor MHC A\*0301, A\*2301, B\*0702, B\*1503

Country United States.

Keywords escape, acute/early infection

References Bernardin et al. 2005

- · Full HIV genomes from nine individuals were analyzed for mutations prior to seroconversion and 7 to 28 days later. It was found that the influence of the host's HLA type was reflected within weeks of infection; a statistically significant number of early nonsynonymous mutations were observed within previously reported CTL epitopes.
- H to Y mutation was observed in position 7.

HXB2 Location Vpr (34–42)

**Author Location** (C consensus)

**Epitope** FPRWLHGL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- FPRWLHGL is an optimal epitope for B\*4201, B\*8101, and B\*0702.

HXB2 Location Vpr (34-42)

Author Location (C consensus)

Epitope FPRWLHGL

Subtype C

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- FPRWLHGL is an optimal epitope for B\*4201, B\*8101, and B\*0702.

HXB2 Location Vpr (34–42)

**Author Location** Vpr (34–42 SF2)

**Epitope** FPRIWLHGL

Epitope name FL9

Immunogen HIV-1 infection

Species (MHC) human (B\*8101)

Keywords acute/early infection

References Altfeld et al. 2001a

- CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- This epitope was recognized by 2/2 individuals expressing B\*8101 allele and 4/8 individuals expressing B\*0702 allele.
- Individuals with long-term nonprogressive and treated chronic HIV-1 infection targeted Vpr more frequently than individuals with treated acute infection.
- Vpr is a frequent target of HIV-1 specific CD8+ T-cells a response was detected in 45% of individuals tested and Vpr and p17 were the most preferentially targeted proteins per unit length by CD8+ T-cells.

HXB2 Location Vpr (34-42)

Author Location Vpr (34-42)

Epitope FPRIWLHGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*8101)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.

 All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vpr (34–42)

**Author Location** Vpr (34–42)

Epitope FPRIWLHGL

**Immunogen** 

Species (MHC) human (B\*8101)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Vpr (34–42)

**Author Location** (C consensus)

**Epitope** FPRWLHGL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*8101)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- FPRWLHGL is an optimal epitope for B\*4201, B\*8101, and B\*0702.

HXB2 Location Vpr (34–42)

Author Location (C consensus)

Epitope FPRPWLHGL

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*8101, B\*4201, B\*0702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords cross-presentation by different HLA, charac-

terizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location Vpr (34–42)

**Author Location** Vpr (34–42)

Epitope FPRIWLHGL

**Epitope name** B7-FL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 1/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 2/4 individuals had detectable responses to this epitope after STI.

HXB2 Location Vpr (34–42)

Author Location Vpr (34–42)

Epitope FPRTWLHGL

Epitope name B7-FL9 Vpr

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** supervised treatment interruptions (STI), es-

cape, early treatment, superinfection

References Altfeld et al. 2002a

- An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.
- The second infecting strain had the variant fpWtwlhgl. The CTL response declined over time and the response to the second variant was lower than to the first one all the time points.

HXB2 Location Vpr (34–42)

Author Location Vpr

Epitope FPRIWLHGL

**Epitope name** FL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A2, B7, B44, Cw5, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.

 One escape mutation, at position 8, FPRIWLHdL was found not to correspond to the most polymorphic residue in the epitope.

HXB2 Location Vpr (34–42)

**Author Location** Vpr (34–42)

Epitope FPRIWLHGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

HXB2 Location Vpr (34–42)

**Author Location** 

Epitope FPRIWLHGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated, traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Vpr (41–49)

**Author Location** Vpr

**Epitope SLGQHIYET** 

Epitope name Vpr41

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: anchored gp120, Vpr Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human, transgenic mouse (A2)

Assay type T-cell Elispot, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced CTL and CD8+ T-cell IFN gamma responses in mice. Responses to the peptide were not detected in 17 HIV+ HLA-A2 subjects.

**HXB2 Location** Vpr (41–57)

Author Location (C consensus)

Epitope GLGQYIYETYGDTWTGV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*66)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location Vpr (46–54)

**Author Location** Vpr (46–54 2001 HIV-1 subtype B cons)

Epitope IYETYGDTW

Epitope name IW9

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (A\*2501)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** escape, immune evasion, optimal epitope, HIV-1

References Liu et al. 2006

• T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.

HXB2 Location Vpr (48–57) Author Location (C consensus) **Epitope** ETYGDTWTGV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6802)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the W7 residue of ETYGDTWTGV are associated with the presence of the HLA presenting molecule in the host.

HXB2 Location Vpr (48–57)

**Author Location** Vpr (48–57)

Epitope ETYGDTWTGV

Immunogen

Species (MHC) human (A\*6802)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an A\*6802 epitope.

HXB2 Location Vpr (48–57)

**Author Location** Vpr (48–57)

**Epitope** ETYGDTWTGV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6802)

Assav type Other

Keywords HLA associated polymorphism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- ETYGDTWTGV was a previously defined A\*6802 presented epitope that encompassed a supertype A\*2 associated polymorphism, YleTYGDTWTGV,in the first position of that epitope.
- The epitope YleTYGDTWTGV is partially embedded in a CTL immunodominant region.

HXB2 Location Vpr (52–62)

**Author Location** Vpr (52–62)

Epitope DTWAGVEAIIR

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Vpr (53–63) Author Location Vpr (53–63) Epitope TWAVEAIIRI Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A1, A3, B7, B14, Cw\*0702, Cw\*0802

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords acute/early infection, early-expressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

HXB2 Location Vpr (55-70)

Author Location Vpr

Epitope AGVEAIIRILQQLLFI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot 40% (28/70) targeted one or more Vpr peptides, and this peptide was the most frequently recognized epitope in Vpr (41%).
- The regulatory proteins Rev and Tat combined contributed to 3%, and the accessory proteins Vif, Vpr and Vpu to 7%, of the total magnitude of HIV-1 specific CTL responses in a subset of 22 HIV-1 infected individuals in whom all HIV-1 proteins were studied.

**HXB2 Location** Vpr (59–67)

Author Location Vpr (58–66 LAI)

Epitope AIIRILQQL

Subtype B

Immunogen

Species (MHC) human (A\*0201)

Keywords optimal epitope

References Altfeld et al. 2001c; Frahm et al. 2007

• C. Brander notes this is an A\*0201 epitope.

HXB2 Location Vpr (59–67)
Author Location Vpr (58–66 SF2)
Epitope AIIRILQQL
Epitope name AL9
Immunogen HIV-1 infection
Species (MHC) human (A\*0201)

**Keywords** acute/early infection **References** Altfeld *et al.* 2001a

- CTL responses against HIV-1 Vpr, Vpu, and Vif were analyzed in multiple HIV-1-infected individuals.
- This epitope was recognized by 8/24 individuals expressing A\*0201 allele.
- Epitope is located within a highly conserved alpha helix in Vpr.
- Individuals with long-term nonprogressive and treated chronic HIV-1 infection targeted Vpr more frequently than individuals with treated acute infection.
- Vpr is a frequent target of HIV-1 specific CD8+ T-cells a response was detected in 45% of individuals tested and Vpr and p17 were the most preferentially targeted proteins per unit length by CD8+ T-cells.
- The A2 epitopes Vpr AIIRLLQQL and p17 SLYNTVATL do not account for the dominance of Vpr and p17, the result holds even when HLA-A2+ individuals are excluded.

HXB2 Location Vpr (59–67)

**Author Location** Vpr (59–)

Epitope AIIRILQQL

Epitope name Vpr-59

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

**Keywords** binding affinity, subtype comparisons, supertype, computational epitope prediction, im-

munodominance

References Altfeld et al. 2001c

- HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.
- Three additional previously described HLA-A2 epitopes were added to the set of 20, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- AIIRILQQL binds to four HLA-A2 supertype alleles: A\*0203, A\*0201, A\*0206 and A\*6802 (highest affinity), but not A\*0202.
- 5/22 individuals with chronic HIV-1 infection recognized this epitope, but with low magnitude responses in ELISPOT.
- 2/12 HLA-A2 patients with acute HIV-1 infection responded strongly to this peptide, but during chronic infection SL9 and Gag-386 tended to be immunodominant while Vpr-59 was weak and sub-dominant.
- One of the the acutely infected individuals, AC13, was HLA A\*0201/68 B44/14 and also had a strong acute response to gp41 epitope SV10 SLLNATDIAV.
- This peptide was shown to be properly processed and presented in TAP-competent B-cell lines *in vitro*.

HXB2 Location Vpr (59–67)
Author Location Vpr (58–66)
Epitope AIIRILQQL
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (A\*0201)

**Keywords** early-expressed proteins **References** Addo *et al.* 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vpr (59–67)
Author Location Vpr (59–67)
Epitope AIIRILQQL
Immunogen HIV-1 infection
Species (MHC) human (A\*0201)

Donor MHC A\*0201, A32, B49, B51, Cw1, Cw7

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** binding affinity, acute/early infection, earlyexpressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

HXB2 Location Vpr (59–67)
Author Location Vpr (59–)
Epitope AIIRILQQL
Epitope name AL9
Immunogen HIV-1 infection
Species (MHC) human (A2)
Keywords acute/early infection
References Goulder et al. 2001a

 Data from patient AC13 suggest a role for this epitope in initial control of viremia in acute infection, as it is one of several subdominant CTL epitopes recognized during the initial decline in viremia.  A CTL response to SL9, SLYNTVATL, was not evident until 18 months post-presentation.

HXB2 Location Vpr (59–67)
Author Location Vpr (59–67 SF2)
Epitope AIIRILQQL
Immunogen HIV-1 infection
Species (MHC) human (A2)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A2+ individuals that had a CTL response to this epitope broken down by group: 1/10 group 1, 0/6 group 2, and 0/4 group 3.

HXB2 Location Vpr (59-67)

**Author Location** 

Epitope AIIRILQQL

Epitope name Vpr-AL9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA A02, 4/35 (11%) recognized this epitope.

HXB2 Location Vpr (59–67)

Author Location Vpr (59-67)

Epitope ALIRILQQL

Epitope name AL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

Keywords escape, optimal epitope

References Allen et al. 2005b

 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For one of the escape variants, a novel CD8 T-cell response equal in magnitude to the wild type, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wild type.  An escape mutation occurred at position 5 of this epitope, alirSlqql.
 Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested,

HXB2 Location Vpr (59-67)

Author Location Vpr

Epitope ALIRILQQL

Epitope name AL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 5, ALIRsLQQL, was found in the most polymorphic residue in the epitope. One escape mutation, at position 3,ALtRILQQL was found not to correspond to the most polymorphic residue in the epitope.

HXB2 Location Vpr (59-67)

**Author Location** Vpr (59–67)

Epitope ALIRILQQL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, HLA binding

Keywords escape, acute/early infection, variant cross-

recognition or cross-neutralization, optimal

epitope

References Altfeld et al. 2005

- The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was recognized both in acute and chronic infection, but slightly more frequently in chronic infection.
- A less common form of this epitope, with the I to L change in the second position, ALIRILQQL binds to HLA-A2 at lower concentrations and can serve as an HLA-A2 epitope during acute infection. It binds well to A\*0201, A\*0202, A\*0203, and A\*0206. This is an example of a less immunogenic form of the epitope, AiIRILQQL becoming the most common circulating form.

HXB2 Location Vpr (59-67)

**Author Location** Vpr (59–67)

Epitope AIIRILQQL

Immunogen HIV-1 infection

Species (MHC) human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind four of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

HXB2 Location Vpr (59-67)

**Author Location** 

Epitope AIIRILQQL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p.17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated, traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Vpr (62–70)

**Author Location** Vpr (62–)

Epitope RILQQLLFI

Epitope name Vpr-62

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** binding affinity, subtype comparisons, supertype, computational epitope prediction

References Altfeld et al. 2001c

• HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences – 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 – 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.

- Three additional previously described HLA-A2 epitopes were added to the set of 20, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- This epitope binds to three HLA-A2 supertype alleles: A\*0202, A\*6802 (strongest affinity) and A\*0203.
- 3/22 chronically infected patients had a weak ELISPOT response to this epitope.
- 0/12 HLA-A2 patients with acute HIV-1 infection responded to this peptide.

HXB2 Location Vpr (62–70) Author Location Vpr (62–70)

 ${\bf Epitope} \ {\tt RILQQLLFI}$ 

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.
   A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector CD8

HXB2 Location Vpr (62–70)

Author Location Vpr

Epitope RILQQLLFI

Epitope name Vpr 62

Subtype M

Immunogen vaccine, in vitro stimulation or selection

Vector/Type: DNA, peptide Adjuvant: In-

complete Freund's Adjuvant (IFA)

**Species (MHC)** human, mouse, humanized mouse (A\*0201)

Assay type Cytokine production, T-cell Elispot

**Keywords** subtype comparisons, computational epitope prediction, escape, TCR usage, variant cross-

recognition or cross-neutralization

References McKinney et al. 2004

- This study examined variant recognition of epitopes presented by A\*0201 and A\*1101. Numerous amino acid substitutions can be introduced into epitopes without disrupting their recognition by CTLs, although epitopes with multiple substitutions were less recognized. An algorithm was constructed for prediction of epitopes capable of inducing responses to a great number of variant epitopes.
- A total of 23 variant forms of Vpr 62 were identified. More than 95% of the variant epitopes were recognized by CTLs from mice immunized with the parental form.
- Vpr 62 epitope (parent or variant form) was present in 96% of HIV sequences of many M group subtypes.

HXB2 Location Vpr (62–70) Author Location Vpr (62–70) Epitope RILQQLLFI Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

 The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized during chronic infection.

HXB2 Location Vpr (62-70)

Author Location Vpr (62-70)

Epitope RILQQLLFI

Immunogen HIV-1 infection

**Species (MHC)** human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind three of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

**HXB2 Location** Vpr

**Author Location** 

**Epitope** 

Immunogen vaccine

Vector/Type: adenovirus HIV component:

Gag-Pol, Nef, Vpr

Species (MHC) mouse

References Muthumani et al. 2002

- Vpr can cause cells to go into G2 arrest, and it surpresses immune cell activation and inflammatory cytokine production, so co-immunization of BALB/c mice with recombinant adenovirus expressing Vpr and HIV-1 antigens Nef or Gag/Pol was tested to see if Vpr reduced the immune response to the other HIV antigens.
- Vpr compromised CD8+ T-cell lytic responses and T-helper proliferative responses in mice co-immunized with Vpr and Nef or Gag/Pol.
- In vitro, Vpr reduced T-cell cytokine production of IL-12 and TNFalpha, indicative of Vpr-mediated immune suppression.

## II-B-18 Tat CTL/CD8 + epitopes

**HXB2 Location** Tat (2–11) **Author Location** (LAI)

Epitope EPVDPRLEPW

Subtype B

Immunogen

**Species (MHC)** (B\*5301)

Keywords optimal epitope

References Addo et al. 2001; Frahm et al. 2007

HXB2 Location Tat (2–11)

**Author Location** 

Epitope EPVDPRLEPW

**Epitope name** Tat-EW10

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*5301)

References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA B\*5301, 3/15 (20%) recognized this epitope.

**HXB2 Location** Tat (2–11)

**Author Location** (C consensus)

Epitope EPVDPNLEPW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** Tat (2–11)

**Author Location** (C consensus)

**Epitope EPVDPNLEPW** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

• EPVDPNLEPW is an optimal epitope.

HXB2 Location Tat (2-11)

Author Location Tat (2–11 BRU)

Epitope EPVDPRLEPW

Epitope name Tat 1

Immunogen HIV-1 infection

Species (MHC) human (B53)

References Addo et al. 2001

- Rev and Tat are expressed early in the virus life cycle and thus may be important targets for CTL against HIV early in infection and for vaccines therefore CTL responses against Tat and Rev were screened using overlapping peptides.
- 11/57 (19.3%) HIV-1 + individuals recognized at least 1 Tat peptide, and 21/57 (37%) responded to at least 1 Rev peptide.
- EPVDPRLEPW was recognized by four individuals, but only two were B53, thus this epitope can probably be presented by other HLA alleles.

HXB2 Location Tat (2-11)

**Author Location** Tat (2–11)

**Epitope** EPVDPRLEPW

Immunogen HIV-1 infection

Species (MHC) human (B53)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Tat (2-11)

**Author Location** Tat

Epitope EPVDPRLEPW

Epitope name EW10

Immunogen HIV-1 infection

Species (MHC) human (B53)

**Assay type** Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords class I down-regulation by Nef

References Bobbitt et al. 2003

• Nef, through Nef-mediated MHC-1 down-regulation, is not the only viral protein to influence levels of HIV-specific CTL recognition. The Rev L60F mutation, a common natural variant, can decrease CTL recognition of late gene products. The Rev mutation impacts the early to late gene switch, reducing late gene product production (Gag, Pol, Env, Vpu, Vpr and Vif), while increasing Nef production, both of which reduce the impact CTL recognition of late gene products. As expected, Rev L60F rendered HIV infected cells more resistant to CTL that recognized epitopes from the late proteins Env and Gag. Gag expression is reduced more than Env, and Gag-specific CTL were more profoundly affected. Conversely CTL against an epitope in an early gene product, Tat, were more effeciently recognized when infected with viruses carrying the Rev L60F mutation.

• Patients in the asymptomic phase with active immune responses had more CTL resistant viruses, with lower Rev activity, lower Gag expression and greater resistance to Gag-specific CTL killing, while viruses isolated from people with AIDS were more sensitive to CTL killing.

HXB2 Location Tat (2-11)

**Author Location** 

Epitope EPVDPRLEPW

Immunogen

Species (MHC) human (B58)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes that this is an B58 epitope.

**HXB2 Location** Tat (2–11)

**Author Location** 

Epitope WPVDPRLEPW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

**HXB2 Location** Tat (3–11)

**Author Location** Tat (3–11 HXB2)

Epitope PVDPRLEPW

Epitope name PW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2501)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assav type CD8 T-cell Elispot - IFNγ

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- This is a newly defined epitope. Positions 5 and 7 in the epitope had potentially experienced positive selection. PVDPRLdPW, PVDPsLEPW and PVDPkLEPW escape variants were found.

HXB2 Location Tat (12–21)

**Author Location** Tat (12–21 SUMA)

Epitope KHPGSQPKTA

Epitope name Tat KA10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A\*1103. A\*2402, B\*1402. B\*1501.

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords dynamics, acute/early infection, characteriz-

ing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTLmediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4 T cell counts through seven years of follow up. In contrast to more rapid progressors, WEAU and BORI, SUMA a broad response to 24 epitopes, with little immunodominance. Two peptides were somewhat more intensely recognized in acute infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

**HXB2 Location** Tat (16–30)

**Author Location** Tat (16–30)

**Epitope** SQPKTACNKCYCKRC

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons

References Novitsky et al. 2002

Keywords escape, immune evasion, optimal epitope, • HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.

- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Tat (17–26) Author Location Tat (17–26)

> **Epitope** QPKTACTTCY Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu
- · All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Tat (17-26)

**Author Location** 

**Epitope** QPKTACTTCY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Tat (20–28)

**Author Location** Tat

Epitope TACNNCYCK

Epitope name 1342

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A03, A23, B49, B57

Country United States. Assay type T-cell Elispot

Keywords binding affinity, computational epitope prediction

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC
- Estimated binding probability for TACNNCYCK: 46%

HXB2 Location Tat (20-29)

**Author Location** Tat

**Epitope** TACNNCYCKK

Epitope name 1279

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A68)

Donor MHC A01, A68, B15, B40, Cw03

Country United States. Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for TACNNCYCKK:74%. This peptide bound A68, not A11.

HXB2 Location Tat (24–32)

**Author Location** Tat (24–32 BORI)

**Epitope** NCYCKKCCY

**Epitope name** Tat NY9

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (A\*2902)

**Donor MHC** A\*2902, B\*1402, Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

**Keywords** dynamics, immunodominance, acute/early infection, characterizing CD8+ T

cells, reversion, viral fitness

References Jones et al. 2004

• Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTLmediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.

- The patient BORI had high viral loads and rapid CD4 decline. BORI mounted 8 detected responses. BORI did not control viral replication well, and escape mutations occurred early and in most epitopes, 6/7 that were precisely identified.
- There were five variants of the NCYCKKCCY epitope in BORI, and new changes kept accruing. kCYCKKCCY was apparent by day 31, kCYCKrCCY by day 218, and kCYCKqCCY by day 556; all conferred escape, the double mutants abrogating the response. NCYCKKyCY and NCYCKKCCc were also transiently present at day 55, but were not tested for CTL escape.

**HXB2 Location** Tat (24–32)

**Author Location** Tat (24–32 WEAU)

Epitope NCYCKRCCF

Epitope name Tat NF9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*2902)

**Donor MHC** A\*2902, B\*4403, B\*0801

Country United States.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, immunodominance, escape, acute/early infection, kinetics, characterizing

CD8+ T cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance. WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.
- There was a weak response to this epitope during acute infection that was lost by early infection. The epitope variant kCYCKRCCF was evident by day 72, and other variants were evident in samples taking at 391 and 772 days, including NCYCKkCCF,tCYCKRCCF, kCYCKsCCF and kCYCKkCCF. It was not determined if these were specifically escape mutations, but the CTL response diminished in vivo as kCYCKRCCF variant came up.

HXB2 Location Tat (29–43)

**Author Location** Tat (29–43)

Epitope KCCFHCQVCFTTKGL

Subtype B

Immunogen HIV-1 infection

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Species (MHC) human
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**Donor MHC** A\*03, A\*24, B\*35, B\*40

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords acute/early infection, variant crossrecognition or cross-neutralization, superinfection, characterizing CD8+ T cells

References Yang et al. 2005b

- An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- A early response to this peptide KCCFHCQVCFTTKGL was detected that waned prior to superinfection. The embedded epitope and HLA presenting molecule were not resolved. The initial and superinfecting strains had different versions of the peptide, oCCFHCQVCFiTKGL and KCClHCQVCFTrKGL respectively.

HXB2 Location Tat (30–37)

**Author Location** Tat (30–37)

**Epitope** CCFHCQVC

Immunogen

Species (MHC) human (Cw\*12)

**Keywords** optimal epitope

**References** Frahm *et al.* 2007

HXB2 Location Tat (30–37)

**Author Location** Tat (30–37)

Epitope CCFHCQVC

Epitope name CC8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*12)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

Keywords escape, optimal epitope

References Allen et al. 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- A mutation occurred at position 3 of this epitope, ccMhcqvc, but significant cross-recognition was observed between the escape variant and the wildtype epitope.

HXB2 Location Tat (30–37)

**Author Location** Tat

Epitope CCFHCQVC

Epitope name CC8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*12)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 3, CClHCQVC, was found in the most polymorphic residue in the epitope.

HXB2 Location Tat (30-37)

**Author Location** Tat

**Epitope** CCFHCQVC

Epitope name CC8

Immunogen

Species (MHC) (Cw\*12)

**Keywords** review, immunodominance, escape, acute/early infection, early-expressed proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

• This review discusses the importance of 3 factors that impact the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides listed as immunodominant in acute HIV-1 infection.

HXB2 Location Tat (30–37)

**Author Location** Tat (30–37)

Epitope CCFHCQVC

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1203)

**Donor MHC** A3, A26, B7, B\*3801, Cw\*0702, Cw\*1203; A\*0201, A\*2501, B18, B51, Cw\*0102,

Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords acute/early infection, early treatment

References Cao et al. 2003

• CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.

- Two individuals recognized this epitope both presented by Cw\*1203.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.

HXB2 Location Tat (30–37)

**Author Location** Tat (30–37)

**Epitope** CCFHCQVC

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1203)

**Donor MHC** A3, A26, B7, B\*3801, Cw\*0702, Cw\*1203;

A\*0201, A\*2501, B18, B51, Cw\*0102,

Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** acute/early infection, early treatment

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- Two individuals recognized this epitope both presented by Cw\*1203.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.

HXB2 Location Tat (30–37)

**Author Location** Tat (30–37 HXB2)

Epitope CCFHCQVC

Epitope name CC8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1203)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- Position 3 in the epitope had potentially experienced positive selection. CClHCQVC and CCFHCQsC escape variants were found.

HXB2 Location Tat (32-41)

**Author Location** Tat (32–41 SUMA)

**Epitope** FHCQVCFMTK

**Epitope name** Tat FK10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

**Keywords** dynamics, epitope processing, immunodominance, escape, acute/early infection, kinetics, characterizing CD8+ T cells, reversion, viral

fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4 T cell counts through seven years of follow up. In contrast to more rapid progressors, WEAU and BORI, SUMA a broad response to 24 epitopes, with little immunodominance. Two peptides were somewhat more intensely recognized in acute infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time. Early in infection, three overlapping epitopes in Tat carried mutations: FHCQVCFMTK, VCFMTKGLGI, and MTKGLGISY. An M->T substitution was evident during acute infection in the first sample, at four days of the onset of symptoms, and a rare second variant was seen at day 20 that added a K->E substitution. The M->T substitution abrogated responses to FHCQVCFtTK, VCFtTKGLGI, but not in the third epitope tTKGLGISY. By day 69 a double mutation was evident that persisted through day 435, F->L and T->K. Variants lHCQVCFMkK, VCFMkKGLGI were not recognized, and impact processing of the MkKGLGISY epitope.

HXB2 Location Tat (36-45)

**Author Location** Tat (36–45 SUMA)

Epitope VCFMTKGLGI

Epitope name Tat VI10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1501)

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

**Keywords** dynamics, epitope processing, immunodominance accepts acute/corly infection kinetics

nance, escape, acute/early infection, kinetics, characterizing CD8+ T cells, reversion, viral

fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4 T cell counts through seven years of follow up. In contrast to more rapid progressors, WEAU and BORI, SUMA a broad response to 24 epitopes, with little immunodominance. Two peptides were somewhat more intensely recognized in acute infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time. Early in infection, three overlapping epitopes in Tat carried mutations: FHCQVCFMTK, VCFMTKGLGI, and MTKGLGISY. An M->T substitution was evident during acute infection in the first sample, at four days of the onset of symptoms, and a rare second variant was seen at day 20 that added a K->E substitution. The M->T substitution abrogated responses to FHCQVCFtTK, VCFtTKGLGI, but not in the third epitope tTKGLGISY. By day 69 a double mutation was evident that persisted through day 435, F->L and T->K. Variants lHCQVCFMkK, VCFMkKGLGI were not recognized, and impact processing of the MkKGLGISY epitope.

HXB2 Location Tat (36-50)

**Author Location** (subtype C)

Epitope VCFQTKGLGISYGRK

Subtype C

Immunogen

Species (MHC) human

Keywords immunodominance, escape

References Novitsky et al. 2001

- This study is provides a survey of CTL responses and full length HIV-1 genome sequences from a C subtype infected Botswanan cohort.
- 17 of 46 patient reacted with Tat immunodominant peptide VCFQTKGLGISYGRK.

Most of the CTL responses occurred despite a mismatch between the autologous viral sequence and peptide – complete matches were seen only in 4 of 19 cases (21%) and the mismatched CTL tended not to respond to the autologous viral peptide indicative of immune escape.

**HXB2 Location** Tat (36–50) **Author Location** Tat (36–50)

Epitope VCFQTKGLGISYGRK

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 8 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Tat (36-52)

Author Location Tat

Epitope VCFTTKALGISYGRKKR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot 28% (19/70) targeted one or more Tat peptides, and this peptide was the most frequently recognized epitope in Tat (27%).
- The regulatory proteins Rev and Tat combined contributed to 3%, and the accessory proteins Vif, Vpr and Vpu to 7%, of the total magnitude of HIV-1 specific CTL responses in a subset of 22 HIV-1 infected individuals in whom all HIV-1 proteins were studied.

**HXB2 Location** Tat (38–47)

**Author Location** (subtype C)

Epitope FQTKGLGISY

Epitope name T38-FY10

Subtype C

Immunogen

Species (MHC) human (B\*1503)

Keywords immunodominance

References Novitsky et al. 2001

- This study is provides a survey of CTL responses and full length HIV-1 genome sequences from a C subtype infected Botswanan cohort.
- 17 of 46 patient reacted with Tat immunodominant peptide VCFQTKGLGISYGRK.
- FQTKGLGISY was the optimal epitope in the peptide VCFQTKGLGISYGRK among B\*1503+ individuals.

**HXB2 Location** Tat (38–47) **Author Location** Tat (38–47)

Epitope FQTKGLGISY

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Tat (38-47)

Author Location (C consensus)

Epitope FQTKGLGISY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location Tat (38-47)

Author Location (C consensus)

**Epitope** FQTKGLGISY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1503)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- FQTKGLGISY is an optimal epitope.

HXB2 Location Tat (39-47)

**Author Location** Tat (39–47 SUMA)

**Epitope MTKGLGISY** 

Epitope name Tat MY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1501)

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** dynamics, epitope processing, immunodominance, escape, acute/early infection, kinetics, characterizing CD8+ T cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4
   T cell counts through seven years of follow up. In contrast
   to more rapid progressors, WEAU and BORI, SUMA a broad
   response to 24 epitopes, with little immunodominance. Two
   peptides were somewhat more intensely recognized in acute
   infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time. Early in infection, three overlapping epitopes in Tat carried mutations: FHCQVCFMTK, VCFMTKGLGI, and MTKGLGISY. An M->T substitution was evident during acute infection in the first sample, at four days of the onset of symptoms, and a rare second variant was seen at day 20 that added a K->E substitution. The M->T substitution abrogated responses to FHCQVCFtTK, VCFtTKGLGI, but not in the third epitope tTKGLGISY. By day 69 a double mutation was evident that persisted through day 435, F->L and T->K. Variants IHCQVCFMkK, VCFMkKGLGI were not recognized, but the CTL response was strong to MkKGLGISY. The authors provide evidence that the F->L and T->K substitutions impact processing of the MTKGLGISY epitope, as the mutations don't abrogate a CTL response to the peptide, but Tat expressed in target cells doesn't allow recognition of the Tat variant.
- MTKGLGISY was the highest level response in acute and early infection.

**HXB2 Location** Tat (39–49)

**Author Location** Tat (38–48)

Epitope ITKGLGISYGR

**Epitope name** Tat-4.8

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

Keywords assay standardization/improvement

References Oxenius et al. 2002a

- This epitope and HLA-A\*6801 presenting molecule were rapidly defined using a modified Elispot assay.
- The 11-mer is the optimal epitope but A\*6801 epitopes tolerate length variation.

HXB2 Location Tat (39-49)

Author Location Tat (39-49)

Epitope ITKGLGISYGR

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Tat (39–49)

**Author Location** Tat (38–48)

Epitope ITKGLGISYGR

Epitope name ITK

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401, Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

006

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot

- IFNγ

Keywords rate of progression, escape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant eptiopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This was one of 8 reactive epitopes found not to vary over time.

HXB2 Location Tat (39–49)

**Author Location** Tat (38–48)

Epitope ITKGLGISYGR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A68)

**Keywords** early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Tat (40–49)

**Author Location** 

Epitope TKALGISYGR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids

that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.

- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

**HXB2 Location** Tat (49–57)

**Author Location** Tat (49–57)

Epitope RKKRRQRRR

Immunogen vaccine

Vector/Type: DNA, DNA with protein boost Strain: B clade LAI HIV component: Gag, Nef, Tat Adjuvant: IL-18

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1

References Billaut-Mulot et al. 2001

- DNA vaccinated BALB/c mice primed and boosted with the multiepitopic vaccine with IL18 showed lymphoproliferative responses 7 weeks post immunization.
- Strong but non-lasting HIV-specific CTL responses were detected by a Cr-release assay and DNA prime/DNA boost was more effective than DNA prime protein boost.
- Immunization with either the multiepitopic DNA or with the mixed DNA vaccine induced HIV-1 specific Th1 cytokines (IL-2 and IFN-gamma)
- Co-administration of IL18 increased T-cell responses but decreased anti-HIV antibody levels.

HXB2 Location Tat (49–57)

**Author Location** Tat (49–57)

Epitope RKKRRQRRR

Immunogen

Species (MHC) mouse

References Kim et al. 1997a

- The Tat peptide RKKRRQRRR when conjugated to a protein can cause that protein to be taken up by APCs and presented to CTL.
- The system was demonstrated by vaccinating mice with an OVA-Tat peptide conjugate and immunizing H-2 K<sup>b</sup> mice.
- The CTL response to the H-2 K<sup>b</sup> specific OVA peptide SIIN-FEKL was stimulated.

 $\textbf{HXB2 Location} \ \ \text{Tat} \ (83 \text{--} 92)$ 

**Author Location** Tat

Epitope GPKESKKKVE

Immunogen

Species (MHC) human (B58)

References De Groot et al. 2001

The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.

- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes could stimulate IFNγ production in an ELISPOT assay.
- GPKESKKKVE was newly identified as an HLA-B58 epitope in this study.

**HXB2 Location** Tat

Author Location Tat

**Epitope** 

Immunogen vaccine

Vector/Type: adeno-associated virus (AAV) HIV component: Env, Rev, Tat Adjuvant:

Species (MHC) mouse (H-2<sup>d</sup>)

References Xin et al. 2001

- An AAV vector expressing HIV-1 env, tat, and rev genes (AAV-HIV vector) was used to vaccinate BALB/c mice.
- A single injection stimulated and long lasting serum IgG, fecal IgA, and HIV-specific CTL.
- Boosting enhanced the humoral response, and IL2 enhanced T-cell immunity.

**HXB2 Location** Tat

Author Location Tat (IIIB)

**Epitope** 

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA), macrophage activating lipopeptide (MALP)

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type T-cell Elispot

References Borsutzky et al. 2003

- BALB/c mice that were vaccinated intranasally with Tat protein plus mucosal adjuvant macrophage-activating lipopeptide-2 (MALP-2) had increased proliferative, antibody, and IFN-gamma producing T-cell responses than did mice with Tat+IFA delivered by the i.p. route.
- IFA as adjuvant stimulated a Th2-dominant response pattern, and MALP-2 as adjuvant shifted to a Th1 response. In animals vaccinated with Tat+MALP-2, IFN-gamma and IL-2 were the most prominent cytokines, with some IL-6. In contrast, in mice vaccinated with Tat+IFA, IL-6 was the dominant cytokine. Secreted IL-4, IL-5 and IL-10 were below the detection limit in both cases.

**HXB2 Location** Tat

**Author Location** Tat

**Epitope** 

Immunogen vaccine

Vector/Type: protein HIV component: Tat Adjuvant: Complete Freund's Adjuvant (CFA), red blood cells

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type Chromium-release assay

Keywords dendritic cells, Th1, Th2, immunotherapy

References Dominici et al. 2003

 BALB/c mice were immunized with Tat protein bound to red blood cells via biotin-avidin conjugation. This antigen delivery system was successfully internalized by dendritic cells, and induced more consistent anti-Tat Abs responses and sightly increased Tat-specific CTL responses relative to Tat with CFA.

HXB2 Location Tat
Author Location Tat
Epitope
Immunogen vaccine

Vector/Type: DNA HIV component: Nef,

Rev, Tat

Species (MHC) human

**Keywords** HAART, ART **References** Calarota *et al.* 1999

- 9/9 HIV-1 + subjects were given one of three DNA vaccinations for nef, rev or tat, and novel proliferative and CTL responses were generated.
- The nef DNA immunization induced the highest and most consistent CTLp activity, IFN-gamma production, and IL-6 and IgG responses.
- Highly active antiretroviral treatment (HAART) did not induce new HIV-specific CTL responses but reduced viral load, while DNA vaccination induced new immune responses but did not reduce viral load – thus this is a potentially complementary and promising combination.

HXB2 Location Tat Author Location Tat Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Froebel *et al.* 1997

- Two HIV-1 infected children with contrasting disease courses were followed longitudinally – one died of AIDS, the other is a long-term non-progressor.
- Reactivity against Gag, Pol, Env and Tat proteins was tested by PBMC bulk cultured cells reacting with protein expressed in vaccinia constructs in autologous EBV transformed B cells.
- The child who progressed consistently had CTL against Pol and Tat.
- The long-term non-progressing child had no detectable CTL, but was heterozygous for a mutation in the CCR5 receptor and for HLA-B49, which has been shown to be associated with slower progression.

HXB2 Location Tat
Author Location Tat
Epitope

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA HIV component: Nef, Rev, Tat Adjuvant: CpG immunostimulatory sequence (ISS)

**Species (MHC)** human **Keywords** review

References Calarota & Wahren 2001

 This review discusses the cellular immune response, and comments on the stimulatory role of CpG motifs and how HIV-1 DNA vaccines can boost the CTL and Th proliferative responses in asymptomatic HIV+ individuals.

HXB2 Location Tat
Author Location Tat
Epitope
Immunogen vaccine

Vector/Type: DNA Strain: B clade BH10 HIV component: Tat Adjuvant: Immune stimulating complexes (ISCOM), CpG immunostimulatory sequence (ISS)

Species (MHC) macaque

References Cafaro et al. 2001

- Macaques (macaca fascicularis) were immunized with HIV-1 Tat on an adenovirus major late promotor in a plasmid with 23 CpG sequences, 12 unmethylated.
- The vaccinated animals contained a primary infection challenge with SHIV89.6P, preventing CD4+ T-cell decline in the animals, suggesting Tat may be useful at blocking viral replication at its early stage.

HXB2 Location Tat
Author Location
Epitope
Subtype B
Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS), responses in children, mother-to-infant transmission

References Aldhous et al. 1994; Kuhn et al. 2002

- Six of nine HIV vertically infected infants had HIV-1 specific CTL responses to vaccinia expressed Tat (4/6), Pol (6/6), Env (1/6), or Gag (1/6), but not all responses were detected at all time points.
- Two of eleven babies that were not infected though born to HIV+ mothers had detectable responses to Tat (1/2), Pol (2/2), Gag (1/2).
- Reviewed in Kuhn et al. [2002].

HXB2 Location Tat Author Location Tat Epitope

Immunogen HIV-1 infection, vaccine

Species (MHC) human

**Keywords** review, escape, early-expressed proteins **References** Gruters *et al.* 2002

- This paper is a review that makes a case for using Tat and Rev as part of a vaccine strategy.
- CTL against Tat and Rev were found preferentially in long term non-progressors.
- Tat/Rev vaccinations of macaques provided protection or reduction in viremia, with high levels of CTL providing protection from challenge, lower levels of CTL having lower viremia, while Gag/Pol vaccinations with did not result in decreased viremia.

• Early expression of Tat/Rev may in part explain the enhanced benefit of a CTL response directed at these proteins, and CTL escape is more prominent in these proteins.

**HXB2 Location** Tat

**Author Location** Tat (BH10)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade BH10 HIV component: Tat Adjuvant: cationic

block copolymer K2

**Species (MHC)** mouse **Donor MHC** H-2d

Assay type proliferation, Chromium-release assay

References Caputo et al. 2003

- Mice were immunized intramuscularly with a plasmid DNA vaccine (HIV-1 pCV-tat DNA) alone or complexed with a cationic block polymer K1, K2, or K5, which block digestion by DNAase I and enhance DNA delivery to APC.
- CTL responses to low dose Tat DNA vaccination with K2 were greatly enhanced relative to responses to DNA alone.

**HXB2 Location** Tat

**Author Location** Tat

**Epitope** 

Immunogen vaccine

Vector/Type: DNA, protein HIV component: Tat Adjuvant: aluminum hydroxide, Ribi adjuvant (MPL+TDM) (RIBI)

Species (MHC) macaque

**Keywords** review, early-expressed proteins **References** Fanales-Belasio *et al.* 2002a

- HIV-1 Tat protein is efficiently taken up by monocyte-derived dendritic cells (MDDCs) and promotes Th1 immune responses.
   A Tat based vaccine can elicit an immune response that can control primary infection in monkeys that are in early stage of in infection with SHIV89.6P.
- Tat-specific CTL activity was detected in four monkeys inoculated with i.m. with pCV-tat.

**HXB2 Location** Tat

**Author Location** 

**Epitope** 

Immunogen in vitro stimulation or selection

Species (MHC)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

Keywords epitope processing, immunodominance, early-

expressed proteins, Th1, adjuvant comparison

References Gavioli et al. 2004

• HIV-1 Tat protein modulates proteasome composition and activity in B and T cells that either express Tat or are treated with exogenous biologically active Tat protein. This results in modification of Ag processing where presentation of immunodominant EBV epitopes is decreased and presentation of subdominant epitopes is increased. The authors suggest that the immunomodulatory effects of endogenous and exogenous Tat may be beneficial in terms of expanding stimulation of responses to subdominant epitopes, and may be useful as an adjuvant.

## II-B-19 Rev CTL/CD8 + epitopes

HXB2 Location Rev (9-23)

Author Location Rev (9–23 HXB2)

Epitope DEELIRTVRLIKLLY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Blazevic et al. 1995

Induces both Th and CTL activities, no HLA restriction analysis performed.

HXB2 Location Rev (11-23)

**Author Location** Rev (14–23)

Epitope KAVRRLIKFLY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

**Keywords** early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Rev (11-23)

**Author Location** Rev (14–23)

**Epitope KAVRRLIKFLY** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5801)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Rev (12-31)

**Author Location** Rev (11–30 SF2)

Epitope LLKAVRLIKFLYQSNPPPNF

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Only one subject had CTL that could recognize vacciniaexpressed LAI Rev.
- This subject had a CTL response to this peptide, and was HLA-A2, A24, B13, B35.

HXB2 Location Rev (14–23)

**Author Location** Rev (14–23 subtype B)

**Epitope** KAVRLIKFLY

Subtype B

Immunogen

Species (MHC) human (B\*5701)

Keywords optimal epitope

References Addo et al. 2001; Frahm et al. 2007

• C. Brander notes this is a B\*5701 epitope.

HXB2 Location Rev (14-23)

**Author Location** Rev (14–23 BRU)

Epitope KAVRIKLFLY Immunogen HIV-1 infection Species (MHC) human (B\*5701)

Keywords cross-presentation by different HLA

References Addo et al. 2001

- Rev and Tat are expressed early in the virus life cycle and thus may be important targets for CTL against HIV early in infection and for vaccines therefore CTL responses against Tat and Rev were screened using overlapping peptides.
- 11/57 (19.3%) HIV-1 + individuals recognized at least 1 Tat peptide, and 21/57 (37%) responded to at least 1 Rev peptide.
- This epitope was also recognized by another individual in whom it was restricted by HLA\*B5801, an allele closely related to HLA\*B5701, suggesting cross-presentation by the two HLA alleles.

**HXB2 Location** Rev (14–23)

**Author Location** Rev (14–23 subtype B)

Epitope KAVRLIKFLY

Subtype B

Immunogen

Species (MHC) human (B\*5801)

Keywords optimal epitope

References Addo et al. 2001; Frahm et al. 2007

• C. Brander notes this is a B\*5801 epitope.

HXB2 Location Rev (14–23)

**Author Location** Rev (14–23 BRU)

Epitope KAVRIKLFLY Immunogen HIV-1 infection Species (MHC) human (B\*5801)

Keywords cross-presentation by different HLA

References Addo et al. 2001

- Rev and Tat are expressed early in the virus life cycle and thus may be important targets for CTL against HIV early in infection and for vaccines therefore CTL responses against Tat and Rev were screened using overlapping peptides.
- 11/57 (19.3%) HIV-1 + individuals recognized at least 1 Tat peptide, and 21/57 (37%) responded to at least 1 Rev peptide.
- This epitope was also recognized by another individual in whom it was restricted by HLA\*B5701, an allele closely related to HLA\*B5801, suggesting cross-presentation by the two HLA alleles.

HXB2 Location Rev (14–23)

**Author Location** Rev

Epitope KTGRLIKLLY

**Epitope name** KY10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 8, ktgrlikLly, was found in the most polymorphic residue in the epitope. One escape mutation, at position 5, ktgrFiklly was found not to correspond to the most polymorphic residue in the epitope.

HXB2 Location Rev (14-23)

**Author Location** 

Epitope KAVRLIKFLY

Immunogen HIV-1 infection

Species (MHC) human (B57, B\*5801)

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

**Keywords** responses in children, mother-to-infant trans-

mission

References Feeney et al. 2005

- Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.
- This epitope was recognized less frequently in children than in adults.

HXB2 Location Rev (14-23)

**Author Location** Rev

Epitope KTVRLIKFLY

Subtype B, C

Immunogen HIV-1 infection

**Species (MHC)** human (B57, B58, B63)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

 HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.

• This epitope was recognized by 20% of B63-positive subjects and 12% of B57/58-positive subjects.

HXB2 Location Rev (14-23)

**Author Location** 

Epitope KAVRLIKFLY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated, traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Rev (14–28)

Author Location (C consensus)

Epitope QAVRIIKILYQSNPY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0205)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression

**References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location Rev (15-23)

**Author Location** Rev (15–23)

**Epitope** TVRLIKFLY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*03)

**Donor MHC** A\*03, A\*24, B\*35, B\*40

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** superinfection, characterizing CD8+ T cells **References** Yang *et al.* 2005b

- An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- The response to this epitope, TVRLIKFLY, was present only after superinfection. The epitope from the first infecting strain had the substitutions tvKlikfly relative to the test peptide, while the second strain shared the sequence TVRLIKFLY.

HXB2 Location Rev (20-28)

**Author Location Rev** 

**Epitope KILYQSNPY** 

Epitope name 1341

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A02, A03, B08, B51, Cw01, Cw07

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for KILYQSNPY: 36%

HXB2 Location Rev (25–39)

**Author Location** Rev (25–39 HXB2)

Epitope SNPPPNPEGTRQARR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Blazevic et al. 1995

Induces both Th and CTL activities, no HLA restriction analysis performed.

HXB2 Location Rev (33-48)

**Author Location** Rev (33–48 HXB2)

Epitope GTRQARRNRRRRWRER

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Blazevic et al. 1995

Induces both Th and CTL activities, no HLA restriction analysis performed.

HXB2 Location Rev (41–56)
Author Location Rev (41–56 HXB2)
Epitope RRRRWRERQRQIHSIS
Subtype B
Immunogen HIV-1 infection
Species (MHC) human

References Blazevic et al. 1995

· Induces both Th and CTL activities.

HXB2 Location Rev (52–60)
Author Location (C consensus)
Epitope IHSISERIL
Subtype C
Immunogen HIV-1 infection
Species (MHC) human (B\*1510)

**Country** South Africa. **Assay type** CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the H2 residue of IHSISERIL are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** Rev (55–63)

**Author Location** Rev

**Epitope ISERILSTY** 

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0101)

**Donor MHC** A\*0101, A\*0301, B\*0801, B\*5101; A\*0101, B\*0801

Country United Kingdom.

**Assay type** CD8 T-cell Elispot - IFNγ, HLA binding **Keywords** escape, acute/early infection, characterizing CD8+ T cells

References Milicic et al. 2005

- CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient. In an early acute infection of the donor, the precise timing of transmission determines the viral variants transmitted.
- The second donor in the study shares A\*0101 and B\*0801 with his partner. The escape variant iserilstF was transmitted, and it abrogates binding to A\*0101.

HXB2 Location Rev (55–63) Author Location Rev (55–63 LAI) Epitope ISERILSTY

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (A1) **Keywords** rate of progression

References van Baalen et al. 1997

- Predicted to be an HLA-A1 epitope based on anchor residues 2S and 9Y.
- Both forms LSGWL(L or I)STY, with intact anchors, were found in an HLA-A1+ individual with Rev-responsive CTL.
- An HLA-A1 individual who did not make a Rev response had lost the C-term anchor, ISGWILS(T or N)S.
- 3/7 long-term non-progressors and 0/5 progressors were positive for HLA-B57 (associated with prolonged survival)
- CTLp frequencies to Rev and Tat were inversely correlated with rapid progression to AIDS, but not Gag, RT or Nef.

**HXB2 Location** Rev (55–63)

**Author Location** Rev (55–63)

**Epitope ISERILSTY** 

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A1)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

HXB2 Location Rev (55–63)

**Author Location** RT Pol (55–63)

**Epitope ISERILSTY** 

Immunogen HIV-1 infection

Species (MHC) human (A1)

Country Spain.

Assay type  $\,$  proliferation, CD8 T-cell Elispot - IFN  $\!\gamma$  , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 8/13 patients recognized this epitope, itw was the most commonly recognized of three A\*01 epitopes tested.

HXB2 Location Rev (56-64)

**Author Location** Rev

Epitope SEWILSTHL

Epitope name SL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 2,SkWILSTHL was found in the most polymorphic residue in the epitope. This is a novel unmapped epitope.

HXB2 Location Rev (57–66)
Author Location Rev (57–66)
Epitope ERILSTYLGR
Immunogen HIV-1 infection
Species (MHC) human (A\*0301)
Keywords optimal epitope
References Frahm et al. 2007

HXB2 Location Rev (57–66) Author Location Rev (57–66) Epitope ERILSTYLGR Epitope name A3-ER10 Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3) Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection.
   1/7 individuals had detectable responses to this epitope after STI.

HXB2 Location Rev (57-66)

**Author Location** 

Epitope ERILSTYLGR

**Subtype** B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

• Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.

- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Rev (58–66)
Author Location Rev (58–66)
Epitope RILSTYLGR
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (A\*0301)
Keywords early-expressed proteins
References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Rev (59–75)

Author Location (C consensus)

Epitope ILSTCLGRPAEPVPLQL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1510)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Rev (66–73) **Author Location** Rev (66–)

Epitope RSAEPVPL

Epitope name Rev66

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Rev Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a low A2-binder, and induced a CTL responses in 1/6 A2 transgenic mice. Responses to the peptide were not detected in 17 HIV+ HLA-A2 subjects.

HXB2 Location Rev (66-75)

**Author Location** 

Epitope RPAEPVPLQL

Epitope name RL10

Immunogen

Species (MHC) human (B7)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B07 epitope.

**HXB2 Location** Rev (66–81)

**Author Location** Rev

Epitope RSAEPVPLQLPPLERL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot 36% (25/70) targeted one or more Rev peptides, and this peptide was the most frequently recognized epitope in Rev (32%).
- The regulatory proteins Rev and Tat combined contributed to 3%, and the accessory proteins Vif, Vpr and Vpu to 7%, of the total magnitude of HIV-1 specific CTL responses in a subset of 22 HIV-1 infected individuals in whom all HIV-1 proteins were studied.

HXB2 Location Rev (66-81)

**Author Location** Rev

Epitope RSAEPAPLQLPPLERL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Novel unmapped epitope.
- RSAEPAPLQLPPLERL shows a mutation over time in position 6, RSAEPvPLQLPPLERL.

HXB2 Location Rev (67-75)

**Author Location (LAI)** 

Epitope SAEPVPLQL

Subtype B

Immunogen

Species (MHC) (B14)

References van Baalen & Gruters 2000

HXB2 Location Rev (67–75)

**Author Location** Rev

Epitope SAEPVPLQL

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords escape

References Schutten et al. 2001

- Molecularly cloned primary NSI macrophage tropic strain 2.1 and SI non-macrophage tropic strain 1.2 were isolated from study participant ACH320 and use to infect irradiated XID mice that had been reconstituted with human PBMC from B14+ seronegative donors – results indicate CTL may favor selective outgrowth of macrophage tropic strains.
- The CTL clone TCC108 specific for SAEPVPLQL, previously described by van Baalen 1997, and van Baalen 1998, was stimulated *in vitro* and given to the mice to apply specific CTL pressure.
- The macrophage-tropic HIV-1 strain #2.1 escaped CTL pressure more efficiently (7/14 animals) than its non-macrophage-tropic counterpart #1.2(SI) the latter isolate was suppressed in 13/14 animals macrophage may serve as a CTL sanctuary and reduced pressure on macrophage tropic HIV strains may allow additional replication to assist with acquisition of escape.
- Specific HIV-1 variants selectively induced by TCC108 were for strain 1.2: SEEPVPLQL, and for strain 2.1: SAEHVPLQL, SAESVPLQL, SVEPVPLQL, SLEPVPLQL, SAEPVPFQL, and SAEPVPFQL.

HXB2 Location Rev (67–75)

**Author Location** Rev (67–75)

Epitope SAEPVPLQL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Keywords** acute/early infection, early-expressed proteins, kinetics

References van Baalen et al. 2002

 Tat, Rev and Nef are the first HIV proteins expressed upon acute infection of T-cells (< 6 hours), and RT is not expressed until after 24 hours. The B14-restricted Rev-SAEPVPLQL specific CD8 T-cell clone TCC108, and the B57-restricted RT-IVLPEKDSW specific CD8 T-cell clone TCL1C11 were co-incubated with CD4+ cultures innoculated with HIV-1 at low MOI. Co-incubation with the Rev-specific CTL resulted in two logs less HIV-1 production in ten days of culture. When the RT epitope was cloned into the Nef gene of the infecting strain, another early expressed protein, it proved as effective as the Rev epitope at inhibiting viral production. A mathematical model of CTL-target interactions suggest early proteins are important for vaccine design.

HXB2 Location Rev (67–75)
Author Location Rev (67–75)
Epitope SAEPVPLQL
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (B14)

Keywords early-expressed proteins

References Addo *et al.* 2002b
CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.

- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Rev (67–75)

Author Location Rev (67–75 IIIB)

Epitope SAEPVPLQL

Immunogen HIV-1 infection

Species (MHC) human (B14, Cw8)

References van Baalen et al. 1998

- The Rev-specific CTL response studied here was from an individual infected with HIV-1 for more than 12 years without developing symptoms Rev and Tat are expressed early and CTL activity against these proteins has been correlated with long-term survival.
- The CTL clone TCC108 specific for this epitope was studied *in vitro*.
- CTLs added immediately after infection suppressed viral production, indicative of CTL interference with viral production prior to lysis CTL-mediated lysis occurred after the onset of progeny viral release, but prior to peak viral production.
- Rapid selection of a E69K mutation, which abolished CTL, recognition was observed.
- The epitope was originally listed as B14, but Cw8 and B14 are in linkage disequilibrium, and in this case were not distinguished (pers. comm., Christian Brander, 1999)

HXB2 Location Rev (67–75)

Author Location (LAI)

Epitope SAEPVPLQL
Subtype B
Immunogen

Species (MHC) human (Cw\*0501)

Keywords optimal epitope

**References** Addo *et al.* 2001; Frahm *et al.* 2007

**HXB2 Location** Rev (67–75) **Author Location** Rev (SF2)

Epitope SAEPVPLQL
Epitope name SL9
Immunogen HIV-1 infection
Species (MHC) human (Cw5)
Keywords acute/early infection
References Goulder et al. 2001a

- Data from patient AC13 suggest a role for this epitope in initial control of viremia in acute infection, as it is one of several subdominant CTL epitopes recognized during the initial decline in viremia.
- A CTL response to SL9, SLYNTVATL, was not evident until 18 months post-presentation.

HXB2 Location Rev (67–75)
Author Location Rev (67–75 SF2)
Epitope SAEPVPLQL
Immunogen HIV-1 infection
Species (MHC) human (Cw5)

**Keywords** HAART, ART, acute/early infection **References** Altfeld *et al.* 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-Cw5+ individuals that had a CTL response to this epitope broken down by group: 2/6 group 1, 0/1 group 2, and 0/2 group 3.

HXB2 Location Rev (67–75)
Author Location Rev (67–75)
Epitope SAEPVPLQL
Immunogen HIV-1 infection
Species (MHC) human (Cw5)

**Donor MHC** A1, A\*0201, B44, B57, Cw5, Cw6

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** binding affinity, acute/early infection, early-expressed proteins

References Cao et al. 2003

• CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.

- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

**HXB2 Location** Rev (67–75) **Author Location** Rev (67–75)

Epitope SAEPVPLQL

Epitope name SL9

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (Cw5)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, Flow cytometric T-cell cytokine

assay

Keywords escape, optimal epitope

References Allen et al. 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- Escape occurred at position 5 of this epitope, saepGplql.

HXB2 Location Rev (67-75)

**Author Location** Rev

Epitope SAEPVPLQL

Epitope name SL9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (Cw5)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 5, SAEPgPLQL, was found not to correspond to the most polymorphic residue in the epitope.

HXB2 Location Rev (67–75)

Author Location Rev (67-75)

Epitope SAEPVPLQL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw5, Cw8)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Rev (67–75)

**Author Location** Rev (69–77 BRU)

Epitope SAEPVPLQL

**Epitope name** Rev SL9

Immunogen HIV-1 infection

Species (MHC) human (Cw8)

**Keywords** HAART, ART, supervised treatment interruptions (STI), acute/early infection

References Addo et al. 2001

- Rev and Tat are expressed early in the virus life cycle and thus may be important targets for CTL against HIV early in infection and for vaccines therefore CTL responses against Tat and Rev were screened using overlapping peptides.
- 11/57 (19.3%) HIV-1 + individuals recognized at least 1 Tat peptide, and 21/57 (37%) responded to at least 1 Rev peptide.
- This epitope is the first HIV-specific CTL epitope resticted by HLA-Cw5.
- This epitope was recognized by 2/5 individuals expressing HLA-Cw8 and by 5/11 individuals expressing Cw5 allele, which differs from Cw8 by 4 amino acids, suggesting promiscuous presentation of the epitope between those HLA molecules.
- Longitudinal data was available for 6 Rev-SL9 responders, who were treated during acute infection, and the response was stable 2 and 12 months after initiaion of HAART, measurements by ELISPOT and flow-based intracellular cytokine staining (ICS) were concordant in two subjects the response was heightened by transient reexposure to antigen with treatment interruption at 12 to 14 months.

HXB2 Location Rev (67–75)

**Author Location** 

Epitope SAEPVPLQL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes,

and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.

- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
- What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Rev (67–75)

Author Location Rev (65-77 BH10, LAI)

Epitope SAEPVPLQL Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is GRSAEPV-PLQLPP) has similarity with transforming growth factor beta binding protein protein I, fragment ARSAEPEVATAPP.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is EPVPLQLPPL) also has similarity with the epidermal growth factor receptor substrate 15, fragment EPVPMSLPPA.

HXB2 Location Rev (67-75)

**Author Location** Rev (67–75)

**Epitope** SAEPVPLQL

Immunogen HIV-1 infection, in vitro stimulation or selec-

Species (MHC) human

Assav type HLA binding

terizing CD8+ T cells

References van Baalen et al. 2005

• A new senstitive, non-readioactive assay, called fluorescent antigen-transfected target cell-CTL (FATT-CTL) assay, was developed to measure antigen-specific cytotoxicity ex vivo. Target cells were generated by nucleofection with DNA vectors encoding antigen-GFP fusion proteins. Flow cytometry was used to quantify viable and dead GFP-positive cells after coculture with different effector:target cell ratios. Cytotoxicity was detected at lower effector:target cell ratios than in standard Cr-release assay. Antigen-specific cytotoxicity was detected ex vivo in PBMCs from HIV-1 infected individuals.

HXB2 Location Rev (72–88)

**Author Location** (C consensus)

Epitope PLQLPPIERLHIDCSES

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*13)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location Rev (73–81)

**Author Location** Rev (73–)

Epitope LQLPPIERL

Epitope name Rev73

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Rev Adjuvant: Incomplete Freund's Adjuvant

(IFA)

**Species (MHC)** human, transgenic mouse (A2)

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced CTL responses in mice. Responses to the peptide were not detected in 17 HIV+ HLA-A2 subjects.

HXB2 Location Rev (75–83)

**Author Location** 

**Epitope** LPPLERLTL

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human

**Keywords** epitope processing, escape

References Yusim et al. 2002

- Keywords assay standardization/improvement, charac- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
  - While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
  - In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.
  - What was learned from proteins where many epitopes have been defined (Gag, Pol, Env and Nef) was used to develop an algorithm to predict where epitopes would be localized in Rev, Tat, Vif, and Vpr. Predictions were made blinded, and then

compared to the first 15 epitopes defined in these proteins; the epitopes were concentrated in the predicted regions.

HXB2 Location Rev (96–104) Author Location Rev (96–) Epitope GMGSPQILV Epitope name Rev96(2M)

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Rev Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (MHC) human, transgenic mouse (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

**References** Corbet *et al.* 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was an intermediate A2 binder, and induced a CD8+ T-cell IFN gamma response in 1/6 mice. Responses were detected in 1/17 HIV+ HLA-A2 subjects.
- The variant gVgspqilv did not elicit a CD8+ T-cell IFN gamma response in transgenic mice, and bound to A2 with low affinity.

HXB2 Location Rev (98-116)

Author Location Rev

Epitope GSTQVSVESPTVLEPGTKE

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A28, A29, B14, B44, Cw8

Country United States.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Novel unmapped epitope. A P->L change occurred in a patient that recognized this peptide, over time: GSTQVSVESITVLEPGTKE

**HXB2 Location** Rev (101–110)

**Author Location** Rev

Epitope QVLGESPTVL

Epitope name QL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Two escape mutations, at positions 5 and 7, QVLGkSPTVL and QVLGEStTVL, were found not to correspond to the most polymorphic residues in the epitope. This is a novel unmapped epitope.

HXB2 Location Rev (102-110)

**Author Location** Rev (102–)

Epitope ILVESPAVL

**Epitope name** Rev102

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Rev Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (MHC) human, transgenic mouse (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder that did not induce CTL or CD8+ T-cell IFN gamma responses in mice, but responses were detected in 2/17 HIV+ HLA-A2 subjects.

HXB2 Location Rev (107–116)

**Author Location** Rev

Epitope PTVLESGTKE

Epitope name 1277

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A68)

**Donor MHC** A11, A68, B42, B45, Cw16, Cw17

Country United States.

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope prediction

References De Groot et al. 2003

Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.

epitope can be presented by A68, but did not bind to A11.

**HXB2 Location** Rev **Author Location Rev Epitope** 

Immunogen vaccine

Vector/Type: DNA with CMV promotor with cationic liposome HIV component: gp160,

Rev

**Species (MHC)** mouse (H-2<sup>d</sup>) References Ishii et al. 1997

- pCMV160/Rev is a DNA vaccine candidate carrying gp160 and Rev linked to a cytomegalovirus (CMV promotor)
- pCMV160/Rev given in conjunction with a cationic liposome gave enhanced DTH, Ab and CTL responses.

**HXB2 Location** Rev **Author Location Rev Epitope** Immunogen vaccine

Vector/Type: DNA HIV component: Rev

Adjuvant: CD40

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1, Th2

References Ihata et al. 1999

• pcRev DNA i.m. vaccination in BALB/c mice induced Th1, Th2 and IgG responses, and enhanced the CTL response to Rev, but did not induce mucosal IgA.

**HXB2 Location** Rev **Author Location** Rev **Epitope** Immunogen vaccine

> Vector/Type: adeno-associated virus (AAV) HIV component: Env, Rev, Tat Adjuvant:

IL-2

**Species (MHC)** mouse (H-2<sup>d</sup>) References Xin et al. 2001

- An AAV vector expressing HIV-1 env, tat, and rev genes (AAV-HIV vector) was used to vaccinate BALB/c mice.
- · A single injection stimulated and long lasting serum IgG, fecal IgA, and HIV-specific CTL.
- Boosting enhanced the humoral response, and IL2 enhanced T-cell immunity.

**HXB2 Location** Rev Author Location Rev

**Epitope** 

Immunogen vaccine

Rev, Tat

Species (MHC) human

Keywords HAART, ART

References Calarota et al. 1999

• 9/9 HIV-1 + subjects were given one of three DNA vaccinations for nef, rev or tat, and novel proliferative and CTL responses were generated.

- Estimated binding probability for PTVLESGTKE:16%. This The nef DNA immunization induced the highest and most consistent CTLp activity, IFN-gamma production, and IL-6 and IgG responses.
  - Highly active antiretroviral treatment (HAART) did not induce new HIV-specific CTL responses but reduced viral load, while DNA vaccination induced new immune responses but did not reduce viral load – thus this is a potentially complementary and promising combination.

**HXB2 Location** Rev

**Author Location** (subtype C)

**Epitope** 

Subtype C

**Immunogen** 

Species (MHC) human

References Novitsky et al. 2001

- This study is provides a survey of CTL responses and full length HIV-1 genome sequences from a C subtype infected Botswanan cohort.
- · Anti-Rev CTL responses were distributed throughout the protein and 27 of 47 subjects (57%) demonstrated HIV-1C Revspecific ELISPOT CTL responses of more than 100 SFC/106 PBMC.

**HXB2 Location** Rev

**Author Location Rev** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA HIV component: Nef, Rev, Tat Adjuvant: CpG immunostimula-

tory sequence (ISS)

Species (MHC) human

Keywords review

References Calarota & Wahren 2001

• This review discusses the cellular immune response, and comments on the stimulatory role of CpG motifs and how HIV-1 DNA vaccines can boost the CTL and Th proliferative responses in asymptomatic HIV+ individuals.

**HXB2 Location** Rev

**Author Location Rev** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Species (MHC) human

Keywords review, escape, early-expressed proteins References Gruters et al. 2002

- This paper is a review that makes a case for using Tat and Rev as part of a vaccine strategy.
- CTL against Tat and Rev were found preferentially in long term non-progressors.
- Vector/Type: DNA HIV component: Nef, Tat/Rev vaccinations of macaques provided protection or reduction in viremia, with high levels of CTL providing protection from challenge, lower levels of CTL having lower viremia, while Gag/Pol vaccinations with did not result in decreased
  - Early expression of Tat/Rev may in part explain the enhanced benefit of a CTL response directed at these proteins, and CTL escape is more prominent in these proteins.

## II-B-20 Vpu CTL/CD8 + epitopes

HXB2 Location Vpu (4–13)

Author Location Vpu

Epitope LVILAIVALV

Immunogen

Species (MHC) human (B7)

References De Groot et al. 2001

- The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.
- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes could stimulate IFNγ production in an ELISPOT assay.
- LVILAIVALV was newly identified as an HLA-B7 epitope in this study using ELISPOT, but could not be shown to bind to B7.

HXB2 Location Vpu (4-13)

Author Location Vpu

Epitope LVILAIVALV

Epitope name 1300

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A03, A24, B07, B38, Cw07, Cw12/13

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for LVILAIVALV: 6%

HXB2 Location Vpu (13-21)

Author Location Vpu (13-)

Epitope VVAAIIAIV

Epitope name Vpu13

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Vpu Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (MHC) human (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

 HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.

This peptide was an intermediate A2 binder that did not induce CTL and CD8+ T-cell IFN gamma responses in mice.
 Responses were detected in 2/17 HIV+ HLA-A2 subjects.

HXB2 Location Vpu (25-40)

Author Location Vpu

Epitope IVFIEYRKLRQRKID

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot – only 2% (2/70) targeted one or more Vpu peptides, including this peptide.
- The regulatory proteins Rev and Tat combined contributed to 3%, and the accessory proteins Vif, Vpr and Vpu to 7%, of the total magnitude of HIV-1 specific CTL responses in a subset of 22 HIV-1 infected individuals in whom all HIV-1 proteins were studied.

HXB2 Location Vpu (29–37)

Author Location Vpu (29–37)

Epitope EYRKILRQR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*3303)

Keywords early-expressed proteins

References Addo et al. 2002b

- CTL responses against regulatory and accessory proteins in 70 HIV-1 infected patients were studied using overlapping peptides and Elispot.
- 28% targeted one or more overlapping Tat peptides; 36%, Rev peptides; 33%, Vif peptides; 40%, Vpr peptides; and 2%, Vpu peptides.
- All known optimally defined epitopes were summarized for the five proteins.

HXB2 Location Vpu (29–37)

**Author Location** Vpu (29–37)

Epitope EYRKILRQR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*3303)

Keywords early-expressed proteins

References Addo et al. 2002a

- Detection of HIV CTL epitopes is rare in Vpu, and this is the first optimally defined Vpu epitope.
- This CTL response was first detected in a long term non-progressor, and 3/6 HLA A\*3303 positive individuals were found to have a CTL response to this epitope.
- HLA A\*3303 is common in West Africa and Asia.

HXB2 Location Vpu (29–37)

**Author Location** Vpu (29–37)

Epitope EYRKILRQR

Immunogen HIV-1 infection

Species (MHC) human (A\*3303) Keywords optimal epitope References Frahm *et al.* 2007

**HXB2 Location** Vpu (62–82) **Author Location** Vpu (65–81)

Epitope AALVEMGHDAPWVVDDL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A2, B7, B44, Cw5, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Novel unmapped epitope. The third position was found to vary over time to AAfVEMGHDAPWVVDDL.

HXB2 Location Vpu (66–82)

**Author Location** (C consensus)

Epitope STMVDMGHLRLLDVNDL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location Vpu (74-82)

Author Location Vpu (74–82 2001 HIV-1 subtype B cons)

Epitope HAPWDVNDL

Epitope name HL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0102)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101, Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** escape, immune evasion, optimal epitope, HIV-1

References Liu et al. 2006

 T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.

• This is a newly defined epitope. Last position (9) in the epitope had potentially experienced positive selection. HAPWDVNDm escape variant was found.

HXB2 Location Vpu

Author Location Vpu

Epitope

Immunogen vaccine

Vector/Type: DNA HIV component: Nef,

Vif, Vpu

**Species (MHC)** mouse (H-2<sup>d</sup>)

**Keywords** subtype comparisons, Th1 **References** Ayyavoo *et al.* 2000

- Splenocytes from BALB/c mice immunized with pVVN-P DNA were incubated with Vif, Vpu or Nef antigens for 3 days and assayed for IL-4 and IFN-gamma levels.
- Antigen stimulation increased IFN-gamma production in pVVN-P immunized mice, indicating a Th1 response.
- IL-4 production was not significantly changed after antigen stimulation compared to control levels.
- Cross-clade CTL activity was also observed: A, B clade, CRF01(AE) clade antigens could serve as targets for the B clade immunization-stimulated CTL an HIV-1 AC recombinant, however, did not stimulate a CTL response, but was expressed at lower levels on the target cell.

## II-B-21 gp160 CTL/CD8 + epitopes

HXB2 Location gp160 (2-10)

**Author Location** gp160 (2–10 IIIB)

Epitope RVKEKYQHL

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*0801 epitope.

**HXB2 Location** gp160 (2–10)

**Author Location** gp160 (2–10 IIIB)

Epitope RVKEKYQHL

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords subtype comparisons

References Sipsas et al. 1997

- HIV IIIB proteins were used to define the range of CTL epitopes recognized by 3 lab workers accidentally infected with HIV-1 IIIB.
- Type-specific epitope, unique to the LAI and IIIB because of a deletion of three amino acids that are present in all other subtype B HIV-1s.
- RVKGIRKNYQHL, a variant found in JRCSF, was not recognized
- This epitope is in the signal sequence of gp120.

**HXB2 Location** gp160 (2–10)

Author Location gp120 (2-10)

Epitope RVKEKYQHL Immunogen HIV-1 infection Species (MHC) human (B8) References Day *et al.* 2001

 B8-restricted CTL accounted for about 1/3 of the total CTL response in one individual.

**HXB2 Location** gp160 (6–12)

Author Location gp120 (6–15 CM243 subtype CRF01)

Epitope TQMNWPNLWK

Epitope name E6-15

Subtype CRF01\_AE

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope after a second stimulation in vitro gave a weak response in HEPS study subject 186 who was HLA A2/A11.

**HXB2 Location** gp160 (6–12)

**Author Location** gp120 (6–15 CM243 subtype CRF01)

Epitope TQMNWPNLWK
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A11)

 ${\bf Keywords} \ \ {\bf subtype} \ comparisons$ 

References Bond *et al.* 2001
• HLA-A11 CRF01 (called subtype E in Bond *et al.*) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.

- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This is one of the new A11 epitopes identified through the streamlined EpiMatrix method, and 2/8 tested FSWs recognized it.
- This epitope was not conserved in other subtypes, and exact matches were rare.

**HXB2 Location** gp160 (30–40)

**Author Location** Env (29–39)

**Epitope** AAENLWVTVYY **Immunogen** HIV-1 infection

Species (MHC) human (B44)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load..
- Less than 2 of 11 patients recognized this epitope.

HXB2 Location gp160 (30-49)

Author Location gp120

Epitope AAEQLWVTVYYGVPVWKEAT

Immunogen HIV-1 infection Species (MHC) human (A11)

Keywords TCR usage

References Weekes et al. 1999b

- Peptide 7035.1: Almost all CD8+ T cells are CD28+ at birth, and the proportion of CD28-CD8+ cells increases with age – this study examines the contribution of CD8+CD28- cells to CTL memory pools for CTL clones specific for two persistent human viruses, CMV and HIV – clones were found to be similarly distributed in the CD28 depleted cell population.
- HIV CTL responses to 3 Env and 2 Gag peptides were studied.
- The clonal composition of the TCR  $V\beta$  responses was studied and was found to be highly focused, with one TCR beta-chain sequence tending to dominate the peptide-specific response clones to this epitope were  $V\beta6$ .

**HXB2 Location** gp160 (30–49)

**Author Location** gp120 (1–20)

**Epitope** ATEKLWVTVYYGVPVWKEAT **Epitope** name ATE

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401,

Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

DO6

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot

- IFNγ

**Keywords** rate of progression, escape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant epitopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This was one of 8 reactive peptides found not to vary over time. It was one of four epitopes that were not precisely defined.

**HXB2 Location** gp160 (31–39)

**Author Location** 

**Epitope** AENLWVTVY

Epitope name AY9

Immunogen

**Species (MHC)** human (B\*1801) **Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*1801 epitope.

HXB2 Location gp160 (31–39)

Author Location gp120 (31–39 HIV-MN)

Epitope AENLWVTVY

Epitope name AY9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*1801)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- This is a newly defined epitope. Position 1 in the epitope had potentially experienced positive selection. AdNL-WVTVY, tdNLWVTVY, tEdLWVTVY, eEdLWVTVY and AEdsWVTVY escape variants were found.

**HXB2 Location** gp160 (31–39)

**Author Location** gp160 (30–38 WEAU)

Epitope AENLWVTVY
Epitope name gp160 AY9
Immunogen HIV-1 infection
Species (MHC) human (B\*4403)

**Donor MHC** A\*2902, B\*4403, B\*0801

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** dynamics, immunodominance, escape, acute/early infection, kinetics, characterizing

CD8+ T cells, reversion, viral fitness

References Jones et al. 2004

• Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.

 The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance. WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.

**HIV CTL/CD8+ Epitope Tables** 

• This was the immunodominant response in acute infection in WEAU, and there was rapid escape in the epitope AENL-WVTVY, with three variants observed by day 30 from the onset of symptoms. Additional mutations continued to develop, so that there were 9 different forms observed through the course of sampling. The variants all conferred different levels of reduction in CTL response, double mutations or anchor mutations tended to cause the greatest reduction: AaNLWVTaY, tNkWVTVY, AgNLWVTVY, AkNLWVTVY, although the double mutant tENLWVTiY elicted a very strong CTL response, suggesting it might not be an escape form.

HXB2 Location gp160 (31–39)

Author Location gp120 (30–38 SF2)

Epitope AENLWVTVY Immunogen HIV-1 infection Species (MHC) human (B44)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B44+ individuals that had a CTL response to this epitope broken down by group: 1/8 group 1, 2/3 group 2, and 3/4 group 3.

**HXB2 Location** gp160 (31–39)

Author Location gp120 (30–38)

**Epitope** AENLWVTVY

Immunogen HIV-1 infection

Species (MHC) human (B44)

References Day et al. 2001

HXB2 Location gp160 (31-39)

Author Location gp120

Epitope AENLWVTVY

Immunogen HIV-1 infection

Species (MHC) human (B44)

**Keywords** epitope processing

References Cao et al. 2002

 AC2 is a B44 restricted CTL clone that recognizes AENL-WVTVY.  CTL could be activated by a fusion protein of an HIV protein and anthrax lethal factor (LFn-HIV) that promotes antigen presenting cell uptake of exogenous protein and allows processing through the MHC class I pathway. This strategy for CTL detection could allow antigen presentation without generation of cells by the standard methods of using live viral vectors carrying a protein, or by loading the cells with peptides and by-passing processing.

**HXB2 Location** gp160 (31–39) **Author Location** (B consensus)

**Epitope** AENLWVTVY

Epitope name AY9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B44)

**Donor MHC** A11, A29, B08, B44, Cw4, Cw7

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

Using a flow-cytometric cytotoxicity assay based on caspase-3
activation in dying target cells, it was shown that the subset of
HIV-1-specific CD8+ T cells secreting both IFN-gamma and
TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
perforin expression was found.

• 1/9 individuals recognized this epitope.

**HXB2 Location** gp160 (31–39)

**Author Location** 

Epitope AENLWVTVY

**Epitope name** AY9

Immunogen

Species (MHC) human (B44)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B44 epitope.

**HXB2 Location** gp160 (31–40)

Author Location gp160 (30–39 WEAU)

Epitope AENLWVTVYY

Immunogen HIV-1 infection

Species (MHC) human (B\*4402)

**Keywords** optimal epitope

**References** Frahm et al. 2007

• C. Brander notes this is a B\*4402 epitope.

**HXB2 Location** gp160 (31–40)

**Author Location** gp160 (30–39 WEAU)

Epitope AENLWVTVYY

Immunogen HIV-1 infection

Species (MHC) human (B44)

Keywords immunodominance, escape

**References** Borrow *et al.* 1997; Borrow & Shaw 1998; Goulder *et al.* 1997a

- Two CTL lines from the patient WEAU were studied one had an optimal peptide of (A)AENLWVTVYY, and the other (A)AENLWVTVY, and both responded equally well with one or two N-term Alanines.
- Rapidly post-infection, a strong immunodominant response was observed against this epitope.
- The naturally occurring forms of the peptide found in WEAU were tested as targets for early WEAU CTLs the form TENL-WVTVY was as reactive as the wild type AENLWVTVY but the forms AKNLWVTVY, AGNLWVTVY, AANLWVTVY did not serve as targets.
- The glutamic acid in the second position is a B44 anchor residue.
- Goulder *et al.* [1997a] and Borrow & Shaw [1998] are reviews of immune escape that summarizes this study in the context of CTL escape to fixation.

**HXB2 Location** gp160 (31–55)

Author Location gp120 (32–56 LAI)

Epitope TEKLWVTVYYGVPVWKEATTTLFCA

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (B18)

References Johnson et al. 1994a

 HLA restricted CTL response to epitope in HIV-1 vaccinia-env vaccinees.

**HXB2 Location** gp160 (31–55)

Author Location gp120 (32-56 LAI)

Epitope TEKLWVTVYYGVPVWKEATTTLFCA

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (B18)

References Ferris et al. 1999; Hammond et al. 1995

• This peptide can be processed for HLA-B18 presentation by both TAP-1/2 independent and dependent pathways.

**HXB2 Location** gp160 (32–40)

**Author Location** Env (92TH023)

**Epitope** DNLWVTVYY **Subtype** B, CRF01\_AE

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost, canarypox, canarypox prime with gp160 boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Gag,

gp120, gp41, Pol

Species (MHC) human (B44)

Country Thailand.

Assay type Chromium-release assay

Keywords vaccine-induced epitopes, variant cross-

recognition or cross-neutralization

References Paris et al. 2004

- 21% (40/187) of Thai adults that recieved ALVAC-HIV with or without gp120 or oligomeric gp160 had a CD8+ T-cell response. HLA-B44 was positively associated with CTL responses, and A33 had a borderline significance association with response. A33/B44/DRB1\*0701 is the most common haplotype in Thailand. B46, present in 30% of the population, was negatively associated with CTL responses, although it did not reach significance. HLA class I serotypes A11, A24, A33, B46 and B75 were the most common found in 245 Thai volunteers.
- 9/11 cases of pCTL activity to Env were in people with B44. The authors suggest some of the response may be directed at the previously mapped B44 Env epitope AENLWVTVYY in HXB2, DNLWVTVYY in their CRF01 ALAVC vaccine 92TH023. B\*4403 is the most common B44 allele among Thais, while B\*4402 is more common among Caucasians; a prior study had shown that B\*4403 may be able to present a broader spectrum of epitopes than B\*4402.

HXB2 Location gp160 (32-40)

**Author Location** gp160 (29–37 SUMA)

**Epitope** ENLWVTVYY Epitope name GP160 EY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A\*1103, A\*2402, B\*1402. B\*1501,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords dynamics, acute/early infection, characterizing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTLmediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4 T cell counts through seven years of follow up. In contrast to more rapid progressors, WEAU and BORI, SUMA a broad response to 24 epitopes, with little immunodominance. Two peptides were somewhat more intensely recognized in acute infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

**HXB2 Location** gp160 (33–42)

Author Location Env (32–41 subtype B)

Epitope KLWVTVYYGV

Subtype B

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade MN HIV component: gp160

Species (MHC) human (A\*0201)

Keywords binding affinity

References Kundu et al. 1998a

- Ten HIV-1 + HLA A2 asymptomatic individuals were given two courses of HIV-1 MN rgp160 vaccine over a 2 year period.
- Two hundred and fifty three HIV-1 peptides of 9 or 10 aa possessing the HLA-A2.1 binding motif (Leu at position 2, Val at the C terminus) were identified in gp160, of which 25 had a high or intermediate binding affinity.
- Eleven peptides were studied that had high HLA-A2 binding affinity - a CTL response was detected to 9/11 peptides in at least 1 individual.
- CTL responses after reimmunization may include recall responses - only individuals with vaccine cross-reactive sequences prior to vaccination showed detectable CTL responses.

**HXB2** Location gp160 (33–42)

Author Location gp120 (32–41 LAI)

Epitope KLWVTVYYGV

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human (A2)

References Dupuis et al. 1995

• CTL from HLA-A2 positive subject react with this peptide.

**HXB2 Location** gp160 (33–42)

**Author Location** Env

Epitope NLWVTVYYGV

Epitope name 1256

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A02, A30, B39

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for NLWVTVYYGV: 84%

**HXB2 Location** gp160 (34–42)

**Author Location** 

Epitope LWVTVYYGV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Assay type Cytokine production, proliferation, Tetra-

mer binding, Intracellular cytokine staining,

Chromium-release assay

References Dagarag et al. 2003

- Telomer length is short in the CD8+ T-cell compartment of HIV-1 infected people, indicating excessive CTL activation and premature senescense. Here human telomerase RT (hTERT) transduction of HIV-1-specific CTL was used to study the functional impact of telomerase. Telomerase expression enhanced proliferative capacity, as well as cytolytic and antiviral capabilities; cytokine production was unchanged. hTERT transduced CTLs were 10-fold more efficient in controlling HIV-1 replication in culture. Thus telomerase transduction can restore CTL mediated cytolysis, and may have therapeutic potential.
- Three polyclonal CD8+ T-cell lines derived from an HIV-1, HLA A\*0201 positive patient were used in this study, including one specific for this epitope.

HXB2 Location gp160 (34-55)

Author Location gp120 (25–46 BRU)

Epitope LWVTVYYGVPVWKEATTTLFCA

Immunogen HIV-1 infection Species (MHC) human (A2)

References Dadaglio et al. 1991

· Defined through peptide blocking of CTL activity, and Env deletions.

**HXB2 Location** gp160 (36–44) **Author Location** Env (35–) Epitope VTVYYGVPV

Epitope name Env35

Immunogen HIV-1 infection, vaccine

*Vector/Type:* peptide *HIV component:* Env Adjuvant: Incomplete Freund's Adjuvant

Species (MHC) human (A2)

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced a CD8+ T-cell IFN gamma response in 1/6 mice. Responses to the peptide were not detected in 17 HIV+ HLA-A2 subjects.

HXB2 Location gp160 (36-46)

Author Location Env

**Epitope** VTVYYGVPVWK

Epitope name Env 47 Subtype M

Immunogen vaccine, in vitro stimulation or selection, computer prediction

Vector/Type: DNA, peptide Adjuvant: In-

complete Freund's Adjuvant (IFA) Species (MHC) human, mouse (A\*1101)

Assay type Cytokine production, T-cell Elispot

**Keywords** subtype comparisons, computational epitope prediction, escape, TCR usage, variant crossrecognition or cross-neutralization

References McKinney et al. 2004

- This study examined variant recognition of epitopes presented by A\*0201 and A\*1101. Numerous amino acid substitutions can be introduced into epitopes without disrupting their recognition by CTLs, although epitopes with multiple substitutions were less recognized. An algorithm was constructed for prediction of epitopes capable of inducing responses to a great number of variant epitopes.
- A total of 9 variant forms of Env 47 were identified. More than 95% of the variant epitopes were recognized by CTLs from mice immunized with the parental form.
- Env 47 epitope (parent or variant form) was present in 82% of HIV sequences of many M group subtypes.

HXB2 Location gp160 (36-46)

**Author Location** gp120 (36–46 CM243 subtype CRF01)

Epitope VTVYYGVPVWR

Epitope name E36-4 Subtype CRF01\_AE

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A11)

Keywords HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers - weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope after a second stimulation in vitro gave a weak response in HEPS study subject 186 who was HLA A2/A11.

**HXB2 Location** gp160 (36–46)

**Author Location** gp120 (36–46 CM243 subtype CRF01)

Epitope VTVYYGVPVWR Subtype CRF01\_AE **Immunogen** HIV-1 infection Species (MHC) human (A11)

**Keywords** subtype comparisons References Bond et al. 2001

- HLA-A11 CRF01 (called subtype E in Bond et al.) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This epitope was not predicted by the EpiMatrix method to be likely to bind to A11, though it served as an epitope in the FSWs, and it was one of the six A11 epitopes that had been previously defined.
- 1/8 tested FSWs recognized this epitope.

• This epitope was only conserved in CRF01 and subtypes B and C, and exact matches were uncommon.

**HXB2 Location** gp160 (36–46)

**Author Location** gp120

Epitope VTVYYGVPVWK
Immunogen HIV-1 infection
Species (MHC) human (A11, A\*6801)

References Threlkeld et al. 1997

- Study of the fine specificity of an A3-like-HLA-super-type epitope (the A3-super-type includes A\*0301, A\*1101, A\*3101, A\*3301, and A\*6801)
- The A3 super-type is characterized as a hydrophobic or hydroxyl containing anchor residue at position 2, and a positive charge in the C-term position.
- While most lines were specific, a promiscuous cloned CTL line was derived from an HIV+ donor that could recognize this epitope presented by either A11 or A\*6801.

**HXB2 Location** gp160 (37–45)

**Author Location** gp160

**Epitope** TVYYGVPVW **Subtype** A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: vaccinia Strain: A clade, B clade, D clade NDK, C clade consensus HIV component: Env

Species (MHC) human

**Donor MHC** A\*2902, A\*2902, B\*1503, B\*1801, Cw\*0202, Cw\*1203; A\*3001, A\*6601, B\*5703, B\*5801, Cw\*0401, Cw\*1801

Country Kenya.

Assay type CD8 T-cell Elispot - IFNγ, Other

Keywords subtype comparisons, variant crossrecognition or cross-neutralization

References McKinnon et al. 2005

- Interclade cross-reactivity of clades A, B, C, and D was tested using recombinant vaccinia-based IFN-gamma Elispot assay. 47/74 women had a positive Elispot response to at least one clade, and cross-clade responses were frequent. Clade A responses were most frequent, as expected, as clade A dominates the Kenyan epidemic (85% of the 47 women responded to clade A Env, 74% to B, 62% to C, and 43% to D). Cross-clade CD8 T-cell responses were common and directed at conserved epitopes.
- There was a greater magnitude of response to A clade peptides in individuals who responded to more than 1 clade; a 2-fold higher response was observed in clade A in 36% (9/25) of these individuals, and the response to A peptides was never lower. TVYYGVPVW responses were detected in 2 women who reacted to all clades tested, A, B, C, and D, and the sequence was identical in all clades.

**HXB2 Location** gp160 (37–46)

Author Location gp120 (37-46 LAI)

**Epitope** TVYYGVPVWK

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component: gp160

Species (MHC) human (A\*0301)

References Johnson et al. 1994b

- Multiple CTL clones obtained from two vaccinees.
- C. Brander notes that this is an A\*0301 epitope in the 1999 database.

**HXB2 Location** gp160 (37–46)

Author Location gp120 (38-41 LAI)

Epitope TVYYGVPVWK

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (A\*0301)

References Johnson et al. 1994a

 Highly conserved epitope recognized by multiple CTL clones from vaccinee.

**HXB2 Location** gp160 (37–46)

Author Location gp120 (37–46 LAI)

Epitope TVYYGVPVWK

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (A\*0301)

References Ferris et al. 1999; Hammond et al. 1995

• This peptide can be processed for HLA-A3.1 presentation by TAP-1/2 independent and dependent pathways.

**HXB2 Location** gp160 (37–46)

Author Location gp120 (37-46 LAI)

Epitope TVYYGVPVWK

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (A\*0301)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0301 epitope.

**HXB2 Location** gp160 (37–46)

Author Location gp120 (37–46 LAI)

**Epitope** TVYYGVPVWK

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (A\*0301)

Keywords acute/early infection

References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.

- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

**HXB2 Location** gp160 (37–46)

Author Location gp120

Epitope TVYYGVPVWK Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

Species (MHC) human (A\*0301)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location gp160 (37–46)

**Author Location** Env

**Epitope** TVYYGVPVWK **Immunogen** vaccine

Vector/Type: DNA

Species (MHC) transgenic mouse (A11)

References Ishioka et al. 1999

 A minigene vaccine construct encoding 6 HLA 2.1 and 3 HLA A11 restricted CTL epitopes, the universal Th cell epitope PADRE (pan-DR epitope) and an ER translocating signal sequence was constructed.

- The epitopes were chosen for dominant recognition by CTLs during HBV and HIV infections in humans.
- HLA transgenic mice were used for quantitating in vivo immunogenicity of DNA vaccines encoding HLA-restricted CTL epitopes.

**HXB2 Location** gp160 (37–46)

Author Location Env

**Epitope** TVYYGVPVWK

Epitope name 1283

Subtype multiple

Immunogen HIV-1 infection

**Species (MHC)** human (A11, A2, A3, A\*6801, B18)

Donor MHC A25, A68, B18, B27; A03, A11, B14, B51,

Cw08, Cw13

Country United States.
Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope prediction, cross-presentation by different HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for TVYYGVPVWK:18% Promiscuous epitope binding to A02, A03, A11, A6801 and B18.

**HXB2 Location** gp160 (37–46)

Author Location gp120 (37-46)

Epitope TVYYGVPVWK

Immunogen vaccine

Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag,

gp120, gp41, Protease

Species (MHC) human (A3)

References Carruth et al. 1999

- The vaccine used was a live recombinant canarypox (CP) virus vaccine containing multiple HIV-1 genes (HIV-1 MN gp120, HIV-1 LAI gp41, HIV-1 LAI Gag, HIV-1 LAI protease)
- CD4+ and CD8+ Gag and Env specific CTL responses were detected in only 1/5 vaccinated volunteers, and were not detectable 1 year after vaccination.
- CTL responses to epitopes SLYNTVATL and TVYYGVPVWK from HIV+ control patients were used as positive controls.
- The study explored why vaccinees were non-responsive non-response was not due to inherent defects or differences in the ability of these individuals to process and present antigen.

**HXB2 Location** gp160 (37–46)

**Author Location** gp120 (37–46 LAI)

Epitope TVYYGVPVWK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords review, escape

References Goulder et al. 1997e; Goulder et al. 1997a

- with the same batch of factor VIII.
- One had a response to this epitope, the other did not. They were tested 6-8 years after infection.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study.

HXB2 Location gp160 (37–46) Author Location gp120 (36-45) Epitope TVYYGVPVWK Immunogen HIV-1 infection Species (MHC) human (A3)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** gp160 (37–46) Author Location gp120 (37–46) Epitope TVYYGVPVWK Immunogen HIV-1 infection Species (MHC) human (A3)

Keywords rate of progression, acute/early infection References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- All patients recognized at least 1 A3 epitope, up to 8 A3 epitopes, but none was clearly dominant.

HXB2 Location gp160 (37-46) **Author Location** 

Epitope TVYYGVPVWK Epitope name Env-VK9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3) References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA A03, 0/20 (0%) recognized this epitope.

**HXB2 Location** gp160 (37–46) Author Location gp160 (37–46) **Epitope** TVYYGVPVWK Immunogen HIV-1 infection

**Species (MHC)** human (A3) **Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4, DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- HLA-identical sibling hemophiliac brothers were both infected Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
  - This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

HXB2 Location gp160 (37–46)

**Author Location** gp120

Epitope TVYYGVPVWK

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A3)

Donor MHC A01, A03, B39, B44, Cw4, Cw6

Assay type T-cell Elispot

Keywords HIV exposed persistently seronegative (HEPS)

References Missale et al. 2004

- HIV-specific T-cell response was tested in patients exposed to blood from a patient with highly replicating HIV; these patients were nosocomially infected with HBV, but uninfected with HIV. HIV-specific T-cell responses were directed to structural and non-structural HIV proteins in 2 patients, suggesting that the virus replicated in these patients sufficiently to prime a cell-mediated immune response that protected them from HIV infection.
- This patient responded to 3/11 HIV epitopes tested in an IFNgamma EliSpot assay. Responses were detected 16 and 20 weeks after exposure, but were lost by week 80.

**HXB2 Location** gp160 (37–46) **Author Location** Env (49–58) Epitope TVYYGVPVWK Immunogen HIV-1 infection **Species (MHC)** human (A3 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location gp160 (37-53) **Author Location** (C consensus)

Epitope TVYYGVPVWKEAKTTLF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*3201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location gp160 (37-53)

Author Location (C consensus)

Epitope TVYYGVPVWKEAKTTLF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*4301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** gp160 (37–53)

**Author Location** (C consensus)

Epitope TVYYGVPVWKEAKTTLF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location gp160 (38-48)

Author Location gp120 (45-55)

Epitope VYYGVPVWKEA

Immunogen HIV-1 infection

Species (MHC) human (Cw7)

References Nehete et al. 1998a

- Three long-term non-progressors and one asymptomatic HIV+ individual were studied and found to have HLA class I Crestricted CD8+ Env-specific CTLs – Cw7 specific CTL were found against three peptides, including this one.
- HLA-C antigens are expressed on lymphoid cells to a lesser extent than either HLA-A or -B.

 HLA-C confers protection against lysis by natural killer cells and by non-MHC-restricted effector T cells and Cw7 directly governs this resistance to lysis – the authors hypothesize that pathogens that inhibit antigen expression and class I expression may particularly down regulate Cw7, thus triggering non-MHC restricted killing.

**HXB2 Location** gp160 (42–51)

Author Location gp120 (42-51 PV22)

Epitope VPVWKEATTT

Immunogen HIV-1 infection

Species (MHC) human (B\*5501)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*5501 epitope.

**HXB2 Location** gp160 (42–51)

Author Location gp120 (42-51 PV22)

**Epitope** VPVWKEATTT

Immunogen HIV-1 infection

Species (MHC) human (B55)

References Brander & Walker 1995

• P. Johnson, unpublished.

**HXB2 Location** gp160 (42–51)

Author Location gp120 (41–55)

Epitope VPVWKEATTT

Immunogen HIV-1 infection

Species (MHC) human (B55)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** gp160 (42–52)

Author Location gp120 (42-52)

**Epitope** VPVWKEATTTL

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*3501 epitope.

**HXB2 Location** gp160 (42–52)

**Author Location** (C consensus)

Epitope VPVWKEAKTTL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

HLA class I restricted CD8+ T-cell responses against HIV-1
were analyzed in African patients. Significantly more responses
were shown to be HLA-B restricted. Viral load, CD4 count,
and thus rate of disease progression were also associated with
HLA-B alleles. In addition, the selection pressure imposed on
HIV-1 by HLA-B alleles was shown to be substantially greater
than by other alleles.

• This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

Author Location (C consensus)
Epitope VPVWKEAKTTL
Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*5301) Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$  Keywords rate of progression, optimal epitope References Kiepiela *et al.* 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses
- were associated with higher viremia.

   VPVWKEAKTTL is an optimal epitope.

HXB2 Location gp160 (42–52) Author Location gp120 (42–52 PV22) Epitope VPVWKEATTTL Immunogen HIV-1 infection Species (MHC) human (B35)

**Keywords** subtype comparisons **References** Cao *et al.* 1997a

- VPVWKEATTTL is the consensus sequence for clades B and D.
- VPVWKDAETTL is the consensus sequence for clade A and it is cross-reactive.
- VPVWKEADTTL is the consensus sequence for clade C and it is cross-reactive.
- VPVWKEADTTL is the consensus sequence for clade E and even with three substitutions still retains some cross-reactivity.

HXB2 Location gp160 (42–52)
Author Location gp120 (41–51)
Epitope VPVWKEATTTL
Immunogen HIV-1 infection
Species (MHC) human (B35)
References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location gp160 (42–52) Author Location Env (41–50) Epitope VPVWKEATTTL Immunogen HIV-1 infection Species (MHC) human (B35) Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay **Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 4/9 patients recognized this epitope.

**HXB2 Location** gp160 (42–52)

Author Location Env (43–52 BH10, LAI)

**Epitope** VPVWKEATTTL **Immunogen** HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this peptide is PVWKEATTTL) has similarity with the platelet membrane glycoprotein IIIA precursor (GLIIIA) (integrin beta-3) (CD61): PLYKEATSTF.

**HXB2 Location** gp160 (42–61) **Author Location** gp120 (49–68)

Epitope VPVWKEATTTLFCASDAKAY

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

 HIV-specific CTL lines developed by ex vivo stimulation with peptide.

**HXB2 Location** gp160 (42–61) **Author Location** gp120 (49–68 SF2)

Epitope VPVWKEATTTLFCASDAKAY

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- Three of these 11 had CTL response to this peptide.
- The responding subjects were HLA-A2, A3, B8, B62; HLA-A3, A24, B7, B38.

**HXB2 Location** gp160 (42–61) **Author Location** gp120 (49–68 SF2)

Epitope VPVWKEATTTLFCASDAKAY

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

• CTL expanded *ex vivo* were later infused into HIV-1 infected patients.

**HXB2 Location** gp160 (42–61) **Author Location** gp120 (11–30)

Epitope VPVWKEATTTLFCASDAKAY

Epitope name VPV

Immunogen HIV-1 infection

Species (MHC) human

Cw\*0702, DR17, DR15, DR51, DR52, DO2,

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ

Keywords rate of progression, escape References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant eptiopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This was one of 8 reactive peptides found not to vary over time. It was one of four epitopes that were not precisely defined.

**HXB2 Location** gp160 (50–59) **Author Location** Env (62–71) **Epitope** TTLFCASDAK Immunogen HIV-1 infection **Species (MHC)** human (A3 supertype) Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

**HXB2 Location** gp160 (51–59) **Author Location** Env (63–71) **Epitope** TLFCASDAK Immunogen HIV-1 infection **Species (MHC)** human (A3 supertype) Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401, • This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

> **HXB2 Location** gp160 (52–61) **Author Location** gp120 (59–68 HXB2) **Epitope** LFCASDAKAY Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*2402)

References Lieberman et al. 1992

- CTL epitope defined by T cell line and peptide mapping.
- C. Brander notes that this is an A\*2402 epitope in the 1999 database.

**HXB2 Location** gp160 (52–61) **Author Location** gp120 (53–62 LAI) **Epitope** LFCASDAKAY Subtype B **Immunogen** HIV-1 infection Species (MHC) human (A\*2402)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is an A\*2402 epitope.

**HXB2 Location** gp160 (52–61) Author Location gp120 (53-62) Epitope LFCASDAKAY

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A24)

Keywords HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

· ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** gp160 (52–61) **Author Location** gp120 (53–62)

**Epitope** LFCASDAKAY Immunogen HIV-1 infection Species (MHC) human (A24)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , T-cell Elispot, CD8 T-cell Elispot granzyme B

**Keywords** characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- Two of seven patients responded to this peptide with GzB producing cells, and a different patient with IFN-gamma producing

**HXB2 Location** gp160 (52–61) **Author Location** gp120 (53–62 LAI) **Epitope** LFCASCAKAY Subtype B

Immunogen HIV-1 infection Species (MHC) human (B38) References Shankar *et al.* 1996

• Uncertain whether optimal, binds A24 as well.

**HXB2 Location** gp160 (52–71) **Author Location** gp120 (59–78)

Epitope LFCASDAKAYDTEVHINVWAT

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

 HIV-specific CTL lines developed by ex vivo stimulation with peptide.

HXB2 Location gp160 (52–71) Author Location gp120 (59–78 SF2)

Epitope LFCASDAKAYDTEVHINVWAT

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-A2 and B-21.

**HXB2 Location** gp160 (59–69)

Author Location (C consensus)

**Epitope** KAYETEVHNVW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- KAYETEVHNVW is an optimal epitope.

HXB2 Location gp160 (59-69)

**Author Location** 

**Epitope** KAYETEVHNVW

Epitope name KW11

Immunogen

Species (MHC) human (B58)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B58 epitope.

HXB2 Location gp160 (61-69)

**Author Location** 

Epitope YETEVHNVW

Epitope name YW9

Immunogen

Species (MHC) human (B\*1801)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*1801 epitope.

HXB2 Location gp160 (61-69)

Author Location gp120 (61–69 HIV-MN)

Epitope YETEVHNVW

Epitope name YW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1801)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101, Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- This is a newly defined epitope. Positions 2 and 5 in the epitope had potentially experienced positive selection. YgTEVH-NVW, YdTEVHNVW, YETEaHNVW and YdTEaHNVW escape variants were found.

**HXB2 Location** gp160 (62–80)

Author Location gp120 (69–88 SF2)

Epitope DTEVHNVWATHACVPTDPN

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-A2 and B-21.

**HXB2 Location** gp160 (64–73)

**Author Location** Env (63–72 SF2)

Epitope EVHNVWATHA

Subtype A, B, C, CRF01\_AE, D

Immunogen HIV-1 infection

Species (MHC) human (A\*2603)

Country Japan.

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay, HLA binding

**Keywords** binding affinity, subtype comparisons, computational epitope prediction, rate of pro-

gression, escape, variant cross-recognition or cross-neutralization

References Kawashima et al. 2005

- A\*26 is associated with slow progression to disease and is common in Asian populations (about 20%). 31/110 HIV peptides that carried the A\*2603 motif ([VTILP] at P2, [ML] at the C-terminus) bound to HLA-A\*2603. Only 2 of these were epitopes and could induce specific CD8 T-cell responses in PBMC from HLA-A\*2603 positive subjects.
- This epitope induced specific CD8+ T cells in chronically infected individuals with A\*2603, but not A\*2601.
- 5 common B clade variants were synthesized. EVHNVWATHA and EVHNiWATHA bound to A\*2603 with equal affinity. Ei-HNVWATHA and EaHNVWATHA bound to A\*2603 with reduced affinity. EmHNVWATHA and EkHNVWATHA could not bind to A\*2603. A CTL clone that recognized EVHN-VWATHA was able kill cells prepulsed with the 3 peptide variants that could bind to A\*2602.
- EVHNVWATHA is the most common form in clades A, B, C, and E (CRF01), but EaHNiWATHA is the most common form in clade D.

**HXB2 Location** gp160 (67–75)

**Author Location** Env (67–)

**Epitope** NIWATHACV **Epitope name** Env67(2I)

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: gp120 Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced CTL and CD8+ T-cell IFN gamma responses in mice. Responses were detected in 1/17 HIV+ HLA-A2 subjects.
- The variant nVwathacv was also immunogenic in transgenic mice, but was not recognized in the 17 people tested.

**HXB2 Location** gp160 (75–84)

Author Location gp120

Epitope VPTDPNPPEV
Immunogen HIV-1 infection
Species (MHC) human (A2)

Assay type Cytokine production, CD8 T-cell Elispot -

IFN $\gamma$ , Tetramer binding

References Höhn et al. 2003

The M. tuberculosis HLA-A2 restricted epitope VLT-DGNPPEV and this HLA-A2 HIV-1 gp120 VPTDPNPPEV epitope are cross-recognized. HLA-A2+ patients with pulmonary tuberculosis exhibit cross-reactivity with the HIV gp160 epitope, and those with HIV-1 infection have cross-reactive responses to M.tuberculosis antigen.

HXB2 Location gp160 (78-86)

Author Location gp120 (77–85)

Epitope DPNPQEVVL

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

References Ogg et al. 1998b

 This epitope was included to illustrate the specificity of HIVtetrameric staining, in a cross-sectional study correlating HLA A\*0201 CTL effector cells and low viral load.

**HXB2 Location** gp160 (78–86)

Author Location gp120 (77–85 SF2)

Epitope DPNPQEVVL

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*3501 epitope.

**HXB2 Location** gp160 (78–86)

Author Location gp120 (77–85 SF2)

Epitope DPNPQEVVL

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

References Tomiyama et al. 1997

- A CTL clone responsive to this epitope was obtained.
- 2/7 B35-positive individuals have a CTL response to this epitope.
- This epitope is highly variable.
- The substitutions: 1N, 3S and 7I, 7L and 9M, 8I, 8K all abrogate specific CTL lysis, while only 8K reduces binding to B\*3501
- The substitution 8V to 8E does not reduce specific CTL activity.

**HXB2 Location** gp160 (78–86)

**Author Location** Env (77–85)

Epitope DPNPQEVVL

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

Keywords HAART, ART

References Ogg et al. 1999

- CTL effector levels were measured after potent ARV therapy using HLA-tetramer complexes for the A\*0201 epitopes SYLVANTVATL and ILKEPVHGV in seven patients, and the B\*3501 epitope DPNPQEVVL in one additional patient.
- Levels of CTL effectors typically decline for 5-7 days and then rebound, fluctuating during the first two weeks of therapy.
- After the early fluctuation, there was a steady exponential decay with a median half-life of 45 days.

**HXB2 Location** gp160 (78–86)

**Author Location** Env (77–85)

Epitope DPNPQEVVL

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Dyer et al. 1999

 CTL specific responses were measured over a 1.3 to 1.5 year period in members of the Sydney Blood Bank Cohort (SBBC) who had been infected with a natural attenuated strain of HIV-1 which was Nef-defective. • Some of these patients had prolonged high levels of CTL effector and memory cells despite low viral load.
• The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with pre-

**HXB2 Location** gp160 (78–86)

**Author Location** 

Epitope DPNPQEVVL Immunogen HIV-1 infection Species (MHC) human (B35) Keywords acute/early infection References Wilson *et al.* 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and a close temporal relationship between the number of circulating HIV-specific T cells and viral load.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

**HXB2 Location** gp160 (78–86)

**Author Location (SF2)** 

Epitope DPNPQEVVL Immunogen HIV-1 infection Species (MHC) human (B35) Keywords rate of progression References Kawana *et al.* 1999

- HLA B35 is associated with rapid disease progression.
- The sequences of 9 previously described HIV-1 B35 CTL epitopes were obtained in 10 HLA B35+ and 19 HLA B35- individuals.
- 3/9 CTL epitopes had substitutions that were more common in B35+ individuals than in B35- individuals, but this was one of the six that had no B35 associated pattern of mutation.

**HXB2 Location** gp160 (78–86)

Author Location gp120 (77–85 SF2)

Epitope DPNPQEVVL Immunogen HIV-1 infection Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

 Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.

- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B35+ individuals that had a CTL response to this epitope broken down by group: 1/2 group 1, 0/2 group 2, and 0/1 group 3.

HXB2 Location gp160 (78-86)

**Author Location** 

Epitope DPNPQEVVL Epitope name Env-DL9 Subtype B

Immunogen HIV-1 infection

**Species** (MHC) human (B35) **References** Sabbaj *et al.* 2003

• Among HIV+ individuals who carried HLA B35, 3/20 (15%) recognized this epitope.

**HXB2 Location** gp160 (78–86)

Author Location gp120 (78-86)

Epitope DPNPQEVVL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

Donor MHC A3, A33, B14, B35, Cw\*0401, Cw\*0802

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** binding affinity, acute/early infection, earlyexpressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

**HXB2 Location** gp160 (78–86) **Author Location** (C consensus)

Epitope DPNPQEMVL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** gp160 (78–86)

Author Location gp120 (47-55)

Epitope DPNPQEVAL

Epitope name DPN

Immunogen HIV-1 infection

Species (MHC) human (B35)

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401,

Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

DQ6

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot

**Keywords** rate of progression, immunodominance, escape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant epitopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This epitope was one of six epitopes found to be under positive selection for escape mutations, and was mostly replaced by an escape variant between days 66 and 369 (dpnpqeAal)and,then replaced by a new escape variant (dpnpqevPl) by day 635.

**HXB2 Location** gp160 (78–86)

Author Location gp120 (77-85 SF2)

Epitope DPNPQEVVL

Immunogen HIV-1 infection

Species (MHC) human (B35, B51)

References Shiga et al. 1996

• Binds HLA-B\*3501 and B\*5101 – binds and kills gp120-vaccinia virus infected cells carrying B35 or B51.

HXB2 Location gp160 (78-86)

Author Location gp120 (77-85)

Epitope DPNPQEVVL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B51)

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** gp160 (78–86)

Author Location gp160 (78–86)

Epitope DPNPQEVVL

Immunogen HIV-1 infection

Species (MHC) human (B51)

**Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

HXB2 Location gp160 (88–96)

**Author Location** gp120 (88–96 HIV-MN)

Epitope NVTENFNMW

Epitope name NW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2501)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

• T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.

• This is a newly defined epitope. Positions 5 and 7 in the epitope had potentially experienced positive selection. NVTEdFdMW. NVTEeFdMW and NVTEsFdMW escape variants were found.

**HXB2 Location** gp160 (89–97)

Author Location gp160

**Epitope** VTEEFNMWK

Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: vaccinia Strain: A clade, B clade, D clade NDK, C clade consensus HIV

component: Env

Species (MHC) human

Donor MHC A\*3201, A\*3601, B\*5301, B\*8101,

Cw\*0401, Cw\*0804; A\*2402, A\*3201, Species (MHC) human (A\*0201) B\*5101, B\*5301, Cw\*0401, Cw\*0602

Country Kenya.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Other

**Keywords** subtype comparisons, crossrecognition or cross-neutralization

References McKinnon et al. 2005

- Interclade cross-reactivity of clades A, B, C, and D was tested using recombinant vaccinia-based IFN-gamma Elispot assay. 47/74 women had a positive Elispot response to at least one clade, and cross-clade responses were frequent. Clade A responses were most frequent, as expected, as clade A dominates the Kenyan epidemic (85% of the 47 women responded to clade A Env, 74% to B, 62% to C, and 43% to D). Cross-clade CD8 T-cell responses were common and directed at conserved
- There was a greater magnitude of response to A clade peptides in individuals who responded to more than 1 clade; a 2-fold higher response was observed in clade A in 36% (9/25) of these individuals, and the response to A peptides was never lower. VTEEFNMWK responses were detected in 2 women who had Env responses to all 4 clades, and clade A gave the highest responses; a VnEEFNMWK variant was in clade B and D, and the clade C Env carried VnEEFNMWe. One woman also reacted with RAIEAQQHL, the other with KNCSFNMTT.
- · Both women that reacted with VTEEFNMWK carried HLA-B\*5301, the only common HLA allele.

HXB2 Location gp160 (89–98)

Author Location Env

**Epitope** VTENFNMWKN

Epitope name 1284 Subtype multiple Immunogen HIV-1 infection

Species (MHC) human (A11, A68 supertype)

**Donor MHC** A01, A68, B15, B40, Cw03; A03, A11, B14,

B51, Cw08, Cw13

Country United States. Assay type T-cell Elispot

Keywords binding affinity, supertype, computational epi-

tope prediction, cross-presentation by differ-

ent HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC
- Estimated binding probability for VTENFNMWKN:17%. This epitope can be presented by the A11, A68 supertype.

**HXB2 Location** gp160 (103–111)

**Author Location** Env (102–110)

Epitope QMHEDIISL

Immunogen HIV-1 infection

Keywords binding affinity, TCR usage

References Kmieciak et al. 1998a

- CTL responses in six patients to four Env epitopes were studied: D2: LLNATAIAV, 5.3: RLRDLLLIV, D1: KLTPLCVTL, and 4.3: QMHEDIISL - all have A2 anchor residues.
- The C terminal epitopes (D2 and 5.3) were highly variable and the variability was considered responsible for limited CTL response, while D1 and 4.3, N-terminal epitopes, were much more conserved and gave evidence of high levels of CTL response in vitro.
- Peptides 4.3 and D1 bound HLA-A\*0201 molecules with high
- Peptides 4.3 and D1 stimulated CTL with a relatively limited TCR  $V\beta$  repertoire.

**HXB2 Location** gp160 (104–112)

**Author Location** gp160 (104–112)

Epitope MHEDIISLW

Immunogen HIV-1 infection

Species (MHC) human (B\*3801)

Keywords optimal epitope References Frahm et al. 2007

HXB2 Location gp160 (104–112)

Author Location gp120 (104-112)

Epitope MHEDIISLW

Immunogen HIV-1 infection

Species (MHC) human (B\*3801)

**Donor MHC** A3, A26, B7, B\*3801, Cw\*0702, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, earlyexpressed proteins

References Cao et al. 2003

• CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFNsecreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.

- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized: 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

**HXB2 Location** gp160 (104–119) Author Location gp120 (111–126 IIIB) Epitope MQEDIISLWDQSLKPC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Macatonia et al. 1991

• Primary CTL response with cells from non-infected donors stimulated by the peptide.

**HXB2 Location** gp160 (105–117) Author Location gp120 (112–124 IIIB) Epitope HEDIISLWDQSLK Immunogen HIV-1 infection Species (MHC) human (A2) References Clerici et al. 1991a

• Helper and cytotoxic T cells can be stimulated by this peptide (T2)

HXB2 Location gp160 (105-117) Author Location gp120 (MN) Epitope HEDIISLWDQSLK Immunogen HIV-1 infection Species (MHC) chimpanzee References Lubeck et al. 1997

- · No epitope-specific CTL were detected in chimpanzees immunized with adenovirus-HIV-1 MN gp160 recombinant despite a response to peptides P18 and T1.
- Helper and cytotoxic T cells have been found to be stimulated by this peptide (T2)

**HXB2 Location** gp160 (105–117) Author Location gp120 (112–124 IIIB) Epitope HEDIISLWDQSLK

Immunogen HIV-1 exposed seronegative

Species (MHC) human

References Pinto et al. 1995

• CTL and T helper cell reactivity in healthcare workers exposed to HIV.

HXB2 Location gp160 (108–116)

**Author Location** Env (107–115 subtype B)

Epitope IISLWDQSL

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human (A\*0201) **Keywords** binding affinity References Kundu et al. 1998a

- two courses of HIV-1 MN rgp160 vaccine over a 2 year period.
- Two hundred and fifty three HIV-1 peptides of 9 or 10 aa possessing the HLA-A2.1 binding motif (Leu at position 2, Val at the C terminus) were identified in gp160, of which 25 had a high or intermediate binding affinity.
- Eleven peptides were studied that had high HLA-A2 binding affinity – a CTL response was detected to 9/11 peptides in at least 1 individual.
- CTL responses after reimmunization may include recall responses - only individuals with vaccine cross-reactive sequences prior to vaccination showed detectable CTL responses.

HXB2 Location gp160 (109–117)

**Author Location** Env (109–117 CM243 subtype CRF01)

Epitope ISLWDQSLK Epitope name E109-117 Subtype CRF01\_AE

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A11)

Keywords HIV exposed persistently seronegative (HEPS)

References Bond et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers - weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not
- This epitope was weakly reactive in the HEPS study subject 265 who was HLA A2/A11, and had been predicted to be a possible A11 epitope using Epimer in Bond et al. [2001]

**HXB2 Location** gp160 (110–118)

**Author Location** Env

Epitope SLWDQSLKP

Epitope name 1328 Subtype multiple Immunogen HIV-1 infection Species (MHC) human (A3)

**Donor MHC** A02, A03, B08, B51, Cw01, Cw07

Country United States. Assav type T-cell Elispot

Keywords binding affinity, computational epitope prediction, immunodominance

References De Groot et al. 2003

- · Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for SLWDQSLKP: 50%.Immunodominant epitope.

HXB2 Location gp160 (112-130) Author Location gp120 (119-139 SF2) Epitope WDQSLKPCVKLTPLCVSLK Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-A2 and B-21.

**HXB2 Location** gp160 (112–131)

Author Location gp120 (MN)

Epitope WDQSLKPCVKLTPLCVTLNC

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A2

Assay type CD8 T-cell Elispot - IFNγ

 $\textbf{Keywords} \ \ \text{assay standardization/improvement, HAART,}$ 

ART

References Chitnis et al. 2003

• 17 perinatally HIV-1 infected children (0.08-16 years) were evaluated for HLA-A2-restricted IFN-gamma CD8+ CTL responses against 4 immunodominant peptides that carry HLA-A2 epitopes. Two peptides were from gp120 (one at position 112, one from the V3 loop), and one each was from gp41 and Gag. 15/17 patients responded to the Gag peptide, 13/17 to the gp41 and the non-V3 gp120 peptides, and 11/17 responded to the V3 loop. 4 children recognized all 4 peptides.

HXB2 Location gp160 (117–126) Author Location Env (72–81) Epitope KPCVKLTPLC Immunogen HIV-1 infection

**Species** (MHC) human (B7) **References** Jin *et al.* 2000b

- This B7 epitope is one of three subdominant CTL responses detected in a long-term non-progressor.
- A dominant B7 epitope was defined using conventional methods, and three additional sub-dominant HLA B7 epitopes were defined by first using a non-anchor based strategy, EpiMatrix, to identify 2078 possible epitopes in the autologous HIV-1, followed by B7 anchor residue prediction to narrow the set to 55 peptides for experimental testing.

HXB2 Location gp160 (117–126)

Author Location Env

**Epitope** KPCVKLTPLC

Epitope name 1295

**Subtype** multiple

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for KPCVKLTPLC: 27%. This
  epitope was previously reported but not confirmed in this study.

HXB2 Location gp160 (121-129)

Author Location Env

Epitope KLTPLCVTL

Immunogen vaccine

Vector/Type: DNA

Species (MHC) transgenic mouse (A\*0201)

References Ishioka et al. 1999

- A minigene vaccine construct encoding 6 HLA 2.1 and 3 HLA A11 restricted CTL epitopes, the universal Th cell epitope PADRE (pan-DR epitope) and an ER translocating signal sequence was constructed.
- The epitopes were chosen for dominant recognition by CTLs during HBV and HIV infections in humans.
- HLA transgenic mice were used for quantitating in vivo immunogenicity of DNA vaccines encoding HLA-restricted CTL epitopes strong responses were observed to all nine epitopes, and CTL memory persisted up to four months after a single injection.

**HXB2 Location** gp160 (121–129)

**Author Location** Env (120–128 subtype B)

Epitope KLTPLCVTL

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human (A\*0201)

**Keywords** binding affinity

References Kundu et al. 1998a

- Ten HIV-1 + HLA A2 asymptomatic individuals were given two courses of HIV-1 MN rgp160 vaccine over a 2 year period.
- Two hundred and fifty three HIV-1 peptides of 9 or 10 aa possessing the HLA-A2.1 binding motif (Leu at position 2, Val at the C terminus) were identified in gp160, of which 25 had a high or intermediate binding affinity.
- Eleven peptides were studied that had high HLA-A2 binding affinity a CTL response was detected to 9/11 peptides in at least 1 individual.
- CTL responses after reimmunization may include recall responses only individuals with vaccine cross-reactive sequences prior to vaccination showed detectable CTL responses.

**HXB2 Location** gp160 (121–129)

Author Location Env (120-128)

Epitope KLTPLCVTL

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** binding affinity, TCR usage **References** Kmieciak *et al.* 1998a

- CTL responses in six patients to four Env epitopes were studied:
   D2: LLNATAIAV, 5.3: RLRDLLLIV, D1: KLTPLCVTL, and
   4.3: QMHEDIISL—all have A2 anchor residues.
- The C terminal epitopes (D2 and 5.3) were highly variable and the variability was considered responsible for limited CTL response, while D1 and 4.3, N-terminal epitopes, were much more conserved and gave evidence of high levels of CTL response *in vitro*.
- Peptides 4.3 and D1 bound HLA-A\*0201 molecules with high affinity.
- Peptides 4.3 and D1 stimulated CTL with a relatively limited TCR  $V\beta$  repertoire.
- In a longitudinal study, the CTL response to the variable D2 epitope diminished over time, while the response to the conserved epitope D1 stayed higher over time.

**HXB2 Location** gp160 (121–129)

**Author Location** Env (134–)

**Epitope** KLTPLCVTL

Epitope name Env-134

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** binding affinity, subtype comparisons, supertype, computational epitope prediction

References Altfeld et al. 2001c

- HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.
- Three additional previously described HLA-A2 epitopes were added to the set of 20, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- 2/22 individuals with chronic HIV-1 infection recognized this epitope in ELISPOT.
- 0/12 acutely infected individuals recognized this epitope.
- KLTPLCVTL binds to four HLA-A2 supertype alleles: A\*0201, A\*0202, A\*0203 and A\*6802 (highest affinity).

HXB2 Location gp160 (121-129)

Author Location Env

462

Epitope KLTPLCVTL

Epitope name Env 134

Immunogen vaccine, in vitro stimulation or selection, com-

puter prediction

Vector/Type: DNA

**Species (MHC)** human, humanized mouse (A\*0201)

Assay type Cytokine production, T-cell Elispot

**Keywords** subtype comparisons, computational epitope prediction, escape, TCR usage, variant cross-

recognition or cross-neutralization

References McKinney et al. 2004

 This study examined variant recognition of epitopes presented by A\*0201 and A\*1101. Numerous amino acid substitutions can be introduced into epitopes without disrupting their recognition by CTLs, although epitopes with multiple substitutions were less recognized. An algorithm was constructed for prediction of epitopes capable of inducing responses to a great number of variant epitopes.

- A total of 19 variant forms of Env 134 were identified of which 10 were recognized by CTLs from transgenic mice immunized with the parental form.
- Env 134 epitope was present in 80% of HIV sequences of diverse M group HIV-1 subtypes.

HXB2 Location gp160 (121-129)

**Author Location** Env

**Epitope KLTPLCVTL** 

Epitope name K9L

Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV component: gp140, gp140ΔV3

**Species (MHC)** transgenic mouse (A\*0201)

**Assay type** Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted
CTL responses against multiple conserved HIV-1 epitopes in
HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw
virus expressing gp140 with deleted V1, V2, and V3 loops
successfully induces neutralizing antibodies against HIV-1. A
live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination
against HIV and measles.

**HXB2 Location** gp160 (121–129)

**Author Location** Env

Epitope KLTPLCVSL

Epitope name P10L

Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV compo-

*nent:* gp140, gp140ΔV3

**Species (MHC)** transgenic mouse (A\*0201)

Assay type Chromium-release assay, Flow cytometric Tcell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted
CTL responses against multiple conserved HIV-1 epitopes in
HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw
virus expressing gp140 with deleted V1, V2, and V3 loops
successfully induces neutralizing antibodies against HIV-1. A
live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination
against HIV and measles.

HXB2 Location gp160 (121–129) Author Location gp120 (120–128 LAI) Epitope KLTPLCVTL Subtype B Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human (A2)

References Dupuis et al. 1995

• CTL from HLA-A2 positive subject react with this peptide.

HXB2 Location gp160 (121–129) Author Location gp120 (120–128) Epitope KLTPLCVTL

Immunogen vaccine

Vector/Type: vaccinia

Species (MHC) human (A2)

References Woodberry et al. 1999

- A polyepitope vaccine was generated in a vaccinia construct that contiguously encoded seven epitopes, all presented by HLA A-2.
- HHD mice have a transgene of HLA A2 linked to the transmembrane and cytotoxic domains of H-2D<sup>d</sup> – this transgene is the only MHC molecule expressed in the mice.
- CTL responses to Gag (77-85) SLYNTVATL, Pol (476-484) ILKEPVHGV, gp120 (120-128) KLTPLCVTL, and Nef (190-198) AFHHVAREL were observed in HIV polytope HHD-vaccinated mice, and these responses were enhanced with vaccinia boost.
- No CTL immune responses were generated against HLA A2restricted HIV epitopes Nef 157-166 (PLTFGWCYKL), Pol 346-354 (VIYQYMDDL), and Nef 180-189 (VLEWRFDSRL)
- Sixteen HLA A2+ patients were tested for their ability to make CTL responses by peptide restimulation in culture with the epitopes selected for inclusion in the polytope – one individual recognized all seven of these epitopes; 7 patients had CTL cultures able to recognize at least one of the epitopes, and 6 of those 7 recognized more than one epitope, but they were not able to test all peptides for all patients; many patients only had three peptides tested.
- KLTPLCVTL was recognized by 3 of the patients.

HXB2 Location gp160 (121–129) Author Location gp120 (120–128) Epitope KLTPLCVTL Immunogen HIV-1 infection Species (MHC) human (A2)

**Keywords** dendritic cells **References** Kundu *et al.* 1998b

- Allogeneic dendritic cells (DCs) were obtained from HLAidentical siblings, pulsed with rgp160 MN or A2-restricted HIV-1 epitope peptides, and infused monthly into six HIVinfected patients.
- 1/6 showed increased env-specific CTL and increased lymphoproliferative responses, 2/6 showed increase only in proliferative responses, and 3/6 showed no change pulsed DCs were well tolerated.
- KLTPLCVTL is a conserved HLA-A2 epitope included in this study all six patients had this sequence as their HIV direct sequence, and a detectable CTL response.
- CTL demonstrated against peptide-coated target, epitope is naturally processed and enhancible with vaccine.

HXB2 Location gp160 (121–129)
Author Location gp120 (120–128)
Epitope KLTPLCVTL
Immunogen HIV-1 infection

Species (MHC) human (A2)

References Kmieciak et al. 1998b

 Increased CTL response to cells expressing a VV construct Δv3 mutant compared with a full-length env gene product.

HXB2 Location gp160 (121–129) Author Location gp120 (121–129) Epitope KLTPLCVSL

Immunogen in vitro stimulation or selection

Species (MHC) human (A2) Keywords dendritic cells References Zarling *et al.* 1999

- This study compares the ability of macrophages and dendritic cells to stimulate primary responses in CD8+ lymphocytes isolated from HLA-appropriate HIV-uninfected donors using peptide-pulsed APC – the dendritic cells performed better as APC for the stimulation of primary responses.
- Strong CTL responses were elicited by the epitopes DRFYK-TLRA and GEIYKRWII when presented by either immature or mature dendritic cells macrophages were not able to prime a CTL response against DRFYKTLRA.
- A weak response to KLTPLCVSL was stimulated using macrophages as the APC.
- No detectable response was observed for the following previously-defined HIV epitopes: KIRLRPGGK, ILKEPVHGV, IRLRPGGK, GPKVKQWPL.

Author Location gp160 (121–129) Author Location gp120 (120–128) Epitope KTLPLCVTL Immunogen HIV-1 infection Species (MHC) human (A2)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** gp160 (121–129) **Author Location** gp120 (121–129 IIIB)

Epitope KLTPLCVTL

Epitope name D1
Subtype B
Immunogen vaccine

Vector/Type: DNA, DNA with protein boost Strain: B clade IIIB HIV component:

gp160, gp160ΔV3 Adjuvant: IL-12

**Species (MHC)** mouse (A2)

Keywords vaccine-specific epitope characteristics

References Kiszka et al. 2002

 Transgenic mice expressing a HLA-A2/Kb chimeric protein were vaccinated with a full length gp160 or with gp160deltaV3, with the V3 loop deleted. Mice given gp160deltaV3 had a broader immune response than those given gp160, with increased responses to conserved HLA-A2 epitopes in the C1 region of gp120, KLTPLCVTL, and the C-term region of gp41, SLLNATAIAV.  Greater resistance was conferred by the gp160deltaV3 than the gp160 vaccine to a challenge of vaccinia expressing heterologous gp160 from primary isolates (VI-06 and 89.6), and the resistance was conferred by CD8+ T-cells.

HXB2 Location gp160 (121-129)

**Author Location** Env (121–)

**Epitope KLTPLCVTL** 

Epitope name Env121

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This epitope was one of the previously identified HLA-A2 epitopes studied.
- 3/17 HIV-infected HLA-A2+ people recognized this epitope.

HXB2 Location gp160 (121–129)

**Author Location** gp160 (121–129)

**Epitope** KLTPLCVTL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

 The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was recognized both during acute and chronic infection, but more often during chronic infection.

**HXB2 Location** gp160 (121–129)

Author Location Env (121–129 HXB2)

Epitope KLTPLCVTL

Epitope name D1

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: multiple epitope immunogen HIV component: p17/p24 Gag,

Pol Adjuvant: IL-12

**Species (MHC)** transgenic mouse (A2)

Assay type Cytokine production, CD8 T-cell Elispot -

IFN $\gamma$ , Chromium-release assay

Keywords vaccine-specific epitope characteristics, vac-

cine antigen design

References Bolesta et al. 2005

- Immunization of transgenic mice with a codon-optimized hGagp17p24-Polp51 DNA plasmid, consisting of clusters of highly conserved CTL epitopes presented by multiple MHC class I alleles, induced 2- to 5-fold higher CD8+ T-cell responses than the corresponding full-length proteins. The modified proteins had the ribosomal frameshift deleted, as well as the potentially immunosuppressive p15, and protease and integrase. This correlated with higher protection against challenge with Gag and Pol expressing recombinant vaccinia virus. Mice immunized with the hGagp17p24-Polp51 also showed an elevated level of type 1 cytokine production as well as an increased titer of p24- and RT-specific IgG2 antibody responses.
- Four A2 gag/pol epitopes were tested, and this Env A2 epitope was used as a negative control.

HXB2 Location gp160 (121-129)

**Author Location** Env (134–142)

Epitope KLTPLCVTL

Immunogen HIV-1 infection

Species (MHC) human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind four of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

HXB2 Location gp160 (155–163)

**Author Location** gp160

**Epitope** KNCSFNMTT

Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: vaccinia Strain: A clade, B clade, D clade NDK, C clade consensus HIV component: Env

Species (MHC) human

**Donor MHC** A\*2402, A\*3201, B\*5101, B\*5301, Cw\*0401, Cw\*1604

Country Kenya.

Assay type CD8 T-cell Elispot - IFNγ, Other

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References McKinnon et al. 2005

• Interclade cross-reactivity of clades A, B, C, and D was tested using recombinant vaccinia-based IFN-gamma Elispot assay. 47/74 women had a positive Elispot response to at least one clade, and cross-clade responses were frequent. Clade A responses were most frequent, as expected, as clade A dominates the Kenyan epidemic (85% of the 47 women responded to clade A Env, 74% to B, 62% to C, and 43% to D). Cross-clade

CD8 T-cell responses were common and directed at conserved epitopes.

• There was a greater magnitude of response to A clade peptides in individuals who responded to more than 1 clade; a 2-fold higher response was observed in clade A in 36% (9/25) of these individuals, and the response to A peptides was never lower. VTEEFNMWK responses were detected in 2 women who had Env responses to all 4 clades, and clade A gave the highest responses; a VnEEFNMWK variant was in clade B and D, and the clade C Env carried VnEEFNMWE. One woman also reacted with RAIEAQQHL, the other one with KNCSFNMTT. KNCSFNMTT was identical in clades A and C, while clade B carried KNCSFNMis, clade D carried KNISFNMTT.

HXB2 Location gp160 (156–165)
Author Location gp120 (156–165)
Epitope NCSFNISTSI
Immunogen HIV-1 infection
Species (MHC) human (Cw\*08)
Keywords epitope processing
References Ferris et al. 1999

- Recognized by CTL clone LWF A5, isolated from a lab worker exposed to HIV-1 in 1985.
- The processing of this epitope is TAP1/2-dependent, as are most Env epitopes, and it contains two N-linked glycosylation sites that are glycosylated in Env.
- Only peptide that has been deglycosylated, a process that changes asparagine (N) to aspartic acid (D) was recognized: the aspartic acid at position 5 was critical, position 1 could be either D or N.
- This peptide also contains a Cys involved in a disulfide linkage but reducing conditions did not effect recognition by CTL clone LWF A5.
- The HIV-1 Env epitopes are typically processed by a TAP1/2 dependent mechanism, which involves cotranslational translocation into the ER, glycosylation, export back into the cytosol, and deglycosylation for processing, and retransport into the ER for the association with class I molecules.
- The particular pathway of generating an epitope may have an impact on the presentation of that epitope, quantitatively as well as qualitatively.

HXB2 Location gp160 (156–165)
Author Location gp120 (156–165 IIIB)
Epitope NCSFNISTSI
Immunogen HIV-1 infection
Species (MHC) human (Cw8)
References Sipsas et al. 1997

- HIV IIIB proteins were used to define the range of CTL epitopes recognized by 3 lab workers accidentally infected with HIV-1 IIIB.
- NCSFNITTSI, a variant found in HIV-1 MN, was not recognized, thus this epitope was type-specific.
- NCSFNISTSI contains two potential N-linked glycosylation sites and cysteine residue, possibly related to the requirement for a high sensitizing dose of peptide for CTL activity.

HXB2 Location gp160 (156–165) Author Location Env (162–171 BH10, LAI) Epitope NCSFNISTSI Immunogen HIV-1 infection Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is STSIRGKVQK) has similarity with the macrophage colony stimulating factor I receptor fragment SISIRLKVQK.

HXB2 Location gp160 (188–207)

Author Location gp120 (193–212 BRU)

Epitope TTSYTLTSCNTSVITQACPK

**Immunogen** HIV-1 infection **Species (MHC)** human (A2)

References Dadaglio et al. 1991

• Defined through blocking CTL activity, and Env deletions.

**HXB2 Location** gp160 (191–200)

Author Location gp120 (194–202 CM243 subtype CRF01)

Epitope YRLINCNTSV
Epitope name E191-200
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A2)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was reactive in HIV+ control study subject 144 who carried HLA-A2.

HXB2 Location gp160 (191-200)

Author Location gp120 (194–202 CM243 subtype CRF01)

Epitope YRLINCNTSV
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A2)
Keywords subtype comparisons

References Bond et al. 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- 1/4 tested FSWs recognized the E clade version of this epitope, which differs from the previously defined B clade version by four amino acids, KLTSCNTSV.
- This epitope was somewhat conserved in 4/8 subtypes: CRF01 (E), B, C, and D.

HXB2 Location gp160 (192–200) Author Location gp120 (192–199) Epitope KLTSCNTSV Epitope name SL9

Immunogen HIV-1 infection Species (MHC) human (A\*02) Keywords HAART, ART References Rinaldo *et al.* 2000

Administration of triple-drug antiretroviral therapy (IDV, 3TC and ZDV) sometimes showed a transient increase and other times failed to increase CTL responses in patients with advanced HIV disease, but there is a stable population of tetramer stained HIV-specific CD8+ CD45RO+ cells that persist after therapy and long periods of virus being below the level of detection.

**HXB2 Location** gp160 (192–200) **Author Location** gp120 (199–207)

Epitope TLTSCNTSV Immunogen HIV-1 infection Species (MHC) human (A\*0201) References Brander *et al.* 1996

- This epitope was recognized by PBMC from 6/14 HIV+ asymptomatic patients.
- This epitope was used along with pol CTL epitope ALQDS-GLEV and a tetanus toxin T helper epitope for a synthetic vaccine.
- This vaccine failed to induce a CTL response, although a helper response was evident.

HXB2 Location gp160 (192–200)

Author Location gp120 (192–199 HXB2R)

Epitope KLTSCNTSV
Immunogen HIV-1 infection
Species (MHC) human (A2)

References Brander et al. 1995

• Epitope predicted on HLA binding motif, and studied in the context of inclusion in a synthetic vaccine.

HXB2 Location gp160 (192–200)
Author Location gp120 (192–199)
Epitope KLTSCNTSV
Immunogen HIV-1 infection
Species (MHC) human (A2)
Keywords HAART, ART

**Keywords** HAART, ART **References** Huang *et al.* 2000

- The single cell ELISPOT assay was optimized and highly specific, and found to work well even after the primary cells had been frozen and thawed.
- Increases in gamma interferon producing cells were observed in response to anti-retroviral therapy using single cell IFNgamma-production ELISPOT.

**HXB2 Location** gp160 (192–200)

**Author Location** gp120 (197–205) **Epitope** TLTSCNTSV

Immunogen peptide-HLA interaction

Species (MHC) human (A2)

References Garboczi et al. 1992

 Crystallization of HLA-A2 molecules complexed with antigenic peptides – refers to Dadaglio et al. 1991.

HXB2 Location gp160 (192-200)

**Author Location** gp120 (161–169)

Epitope ILRSCNTSV

Epitope name ILR

Immunogen HIV-1 infection Species (MHC) human (A2)

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401, Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, escape **References** Oxenius *et al.* 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant epitopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This was one of 8 reactive epitopes found not to vary over time.

HXB2 Location gp160 (192-211)

**Author Location** gp120 (199–219 SF2)

Epitope SLTSCNTSVITQACPKVSFE

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-A2, -B21.

**HXB2 Location** gp160 (199–207)

**Author Location** Env (202–210)

Epitope SVITQACPK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*1101)

Keywords subtype comparisons, TCR usage

References Fukada et al. 2002

- Counterparts for eight known clade B HLA A\*1101 epitopes were generated for clade E (CRF01). Three epitopes, identical among clade A-E, were cross-reactive and recognized by clade E infected individuals. The clade E and B analogs to three more HLA A\*1101 epitopes was recognized in a clade specific manner. Two other HLA A\*1101 clade B defined epitopes were found not to have stimulated a response in clade E infected individuals.
- SVITQACPK was found to elicit clade-specific responses in clade B (SVITQACPK is most common, sAitqacpk is most common variant in clade A, C and D) and clade E (saiKqacpk is most common). SVITQACPK was recognized by CTL from 3/5 B clade infected Japanese subjects, and aiKqacpk by CTL

B clade exclusive epitope.

• The binding of the three variant peptides to HLA A\*1101 was comparable, implicating TCR interaction differences.

HXB2 Location gp160 (199-207) Author Location gp160 (199-207) Epitope SVITQACPK Immunogen HIV-1 infection Species (MHC) human (A\*1101) Keywords optimal epitope References Frahm et al. 2007

HXB2 Location gp160 (201-225) Author Location gp120 (201–225 LAI)

Epitope ITQACPKVSFEPIPHYCAPAGFAI

Subtype B Immunogen vaccine

> Vector/Type: vaccinia HIV component: gp160

Species (MHC) human (CD4+ CTL)

References Johnson et al. 1994b; Johnson et al. 1994a

• CD4+ CTL isolated from LAI IIIB gp160 vaccinees.

HXB2 Location gp160 (202-221) Author Location gp120 (209-228)

Epitope TQACPKVSFEPIPIHYCAPA

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

• HIV-specific CTL lines developed by ex vivo stimulation with peptide.

HXB2 Location gp160 (202-221)

Author Location gp120

Epitope TQACPKVSFEPIPIHYCAPA

Immunogen HIV-1 infection

Species (MHC) human

Keywords TCR usage

References Weekes et al. 1999b

- Peptide 740.18: Almost all CD8+ T cells are CD28+ at birth, and the proportion of CD28-CD8+ cells increases with age this study examines the contribution of CD8+CD28- cells to CTL memory pools for CTL clones specific for two persistent human viruses, CMV and HIV - clones were found to be similarly distributed the CD28 depleted cell population.
- HIV CTL responses to 3 Env and 2 Gag peptides were studied.
- The clonal composition of the TCR  $V\beta$  responses were studied and was found to be highly focused, with one TCR beta-chain sequence tending to dominate the peptide-specific response clones to this epitope were  $V\beta 13.1$ .

HXB2 Location gp160 (202-221)

Author Location gp120

Epitope TQACPKVSFEPIPIHYCAPA

Immunogen HIV-1 infection

Species (MHC) human

References Weekes et al. 1999a

from 0/7 E clade infected Thai subjects, so this seems to be a • Peptide 740.18: Memory CTL specific for HIV-1 may contribute to oligoclonal expansions within the CD57+ CD28- CD8+ CTLp populations.

HXB2 Location gp160 (202-221)

Author Location gp120 (209–228 SF2)

Epitope TQACPKVSFEPIPIHYCAPA

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1
- · Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- One of these 11 had CTL response to this peptide.

**HXB2 Location** gp160 (202–221)

**Author Location** gp120 (209–228 SF2)

Epitope TQACPKVSFEPIPIHYCAPA

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

• CTL expanded ex vivo were later infused into HIV-1 infected patients.

HXB2 Location gp160 (207-216)

Author Location gp120 (subtype A)

Epitope KMTFEPIPIH

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (A29)

Keywords subtype comparisons

References Cao et al. 2000

- HIV-1 subtypes A and D dominate the Ugandan epidemic, and a vaccine trial using B clade antigen is underway – this study addresses relative levels of cross-reactive CTL responses in Ugandans to A, D, and B clade recombinant vaccinia viruses expressing Gag, Env, Pol, RT or Nef from HIV-1 clades A, B, and D.
- Proteins corresponding to the subtype of the infecting strains tended to trigger higher levels of CTL response measured by percent specific lysis, but there was extensive inter-subtype cross-reactivity with B clade proteins and the co-circulating
- CTL derived from subtype A clade infection (patient SP 528), recognized the subtype A version of the peptide (KMS-FEPIPIH), had a slightly reduced specific lysis using the B clade version of the peptide (KVSFEPIPIH), and no lysis using the D clade version of the epitope (KVTFEPIPIH)
- Patient SP 528 is HLA A1, A29, B57, B81, Bw4, Bw6.

**HXB2 Location** gp160 (207–224)

**Author Location** (C consensus)

Epitope KVSFDPIPIHYCAPAGYA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0401)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location gp160 (208-216)

**Author Location** Env

Epitope VSFEPIPIH

Epitope name 1329

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A03, A23, B49, B57; A03, A24, B27, B57,

Cw13, Cw18

Country United States.

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for VSFEPIPIH: 58%

**HXB2 Location** gp160 (208–217)

Author Location gp120 (subtype B)

Epitope VSFEPIPIHY

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A29)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix systemic CD8+ T cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

**HXB2 Location** gp160 (208–217)

Author Location gp120 (263–272)

Epitope VSFEPIPIHY

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A29)

Keywords HIV exposed persistently seronegative

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers. HXB2 Location gp160 (208–217)

Author Location gp120

**Epitope** VSFEPIPIHY

Immunogen HIV-1 infection

Species (MHC) human (A29)

Assay type Intracellular cytokine staining

Keywords immunodominance, genital and mucosal im-

munity

References Kaul et al. 2003

- Predefined immunodominant peptide responses were used to compare CD8+ T cells responses in the blood and cervix of 16 HIV+ Kenyan sex workers. Cervical responses were detected in 8/10 women from whom adequate samples could be obtained. The frequency of the CD8+ T cell response in the genital tract was comparable to the blood, with a trend toward being slightly higher.
- The immunodominant response was to this epitope in the PBMC of 1/16 patients (Kaul *et al.* 2001, AIDS, 107:1303).

HXB2 Location gp160 (208-219)

**Author Location** Env

Epitope VSFEPIPPHYCA

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords epitope processing

References Cao et al. 2002

- SP 511 is an A2 restricted CTL clone generated from a Ugandan subject that recognizes VSFEPIPPHYCA.
- CTL could be activated by a fusion protein of an HIV protein and anthrax lethal factor (LFn-HIV) that promotes antigen presenting cell uptake of exogenous protein and allows processing through the MHC class I pathway. This strategy for CTL detection could allow antigen presentation without generation of cells by the standard methods of using live viral vectors carrying a protein, or by loading the cells with peptides and by-passing processing.

**HXB2 Location** gp160 (208–219)

**Author Location** Env

Epitope VSFEPIPPHYCA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A\*0202, A\*0301, B\*0702, B\*1516

Country United States.

Keywords escape, acute/early infection

References Bernardin et al. 2005

- Full HIV genomes from nine individuals were analyzed for mutations prior to seroconversion and 7 to 28 days later. It was found that the influence of the host's HLA type was reflected within weeks of infection; a statistically significant number of early nonsynonymous mutations were observed within previously reported CTL epitopes.
- K to E mutation was observed in position 4.

HXB2 Location gp160 (209–217)

**Author Location** (C consensus)

**Epitope** SFDPIPIHY **Subtype** C

Immunogen HIV-1 infection

Species (MHC) human (A\*29)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the S1 residue of SFDPIPIHY are associated with the presence of the HLA presenting molecule in the host

HXB2 Location gp160 (209-217)

**Author Location** (LAI)

**Epitope SFEPIPIHY** 

Subtype B

Immunogen

Species (MHC) human (A\*2902)

Keywords optimal epitope

References Altfeld 2000; Frahm et al. 2007

HXB2 Location gp160 (209–217)

Author Location gp160 (207–215 BORI, WEAU)

Epitope SFEPIPIHY

Epitope name gp160 SY9

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (A\*2902)

**Donor MHC** A\*2902, B\*1402, Cw\*0802; A\*2902

B\*4403, B\*0801

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

**Keywords** dynamics, immunodominance, escape

acute/early infection, characterizing CD8+ T

cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined, WEAU and BORI, had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape. This epitope was recognized in both patients.
- The patient BORI had high viral loads and rapid CD4 decline. BORI mounted 8 detected responses. BORI did not control viral replication well, and escape mutations occurred early and in most epitopes, 6/7 that were precisely identified. The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance. WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.

- Four escape variants to the SFEPIPIHY epitope were found in the patient BORI. SFdPIPIHY came up first, at day 55 from onset of symptoms, and caused a reduced cytotoxic response. By day 218, two rare forms were found, SIEPIPIHf and SiEPIPIHf. By day 556, only tFEPIPIHY was found. The weakest response was detected in the double mutant, SiEPIPIHf, yet tFEPIPIHY was the form that persisted.
- In WEAU, a minor variant, SsEPIPIHY was present at day 41.
   The SIEPIPIHf variant first came up day 136, gave a reduced CTL response, and then came to be the dominant form. Other variants were SFEPIPINY and SFEPIPIdf.

**HXB2 Location** gp160 (209–217)

Author Location gp120 (213-221 SF2)

Epitope SFEPIPIHY
Immunogen HIV-1 infection
Species (MHC) human (A29)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- A\*2902, Previously described and newly defined optimal epitopes were tested for CTL response.
  - Number of HLA-A29+ individuals that had a CTL response to this epitope broken down by group: 1/3 group 1, 0/0 group 2, and 0/0 group 3.

**HXB2 Location** gp160 (209–217)

**Author Location** gp120 (209–217)

Epitope SFEPIPIHY

Immunogen HIV-1 infection

Species (MHC) human (A29)

**Donor MHC** 1261: A\*0201, A29, B58, B62, Cw\*0304,

Cw\*1601; 1168: A\*0201, A29, B44, B60,

Cw3, Cw16

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

• All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef. • 1/9 individuals recognized this epitope Tat, Vpr. and Env.

- Two subjects recognized this epitope during primary infection, both in the context of A29.
- · All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

HXB2 Location gp160 (209-217)

Author Location (C consensus)

**Epitope** SFDPIPIHY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A29)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location gp160 (209-217)

Author Location (B consensus)

**Epitope SFEPIPIHY** 

Epitope name SY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A29)

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

• Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.

HXB2 Location gp160 (212-231)

Author Location gp120

Epitope PIPIHYCAPAGFAILKCNNK

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords TCR usage

References Weekes et al. 1999b

- Peptide 740.19: Almost all CD8+ T cells are CD28+ at birth, and the proportion of CD28-CD8+ cells increases with age this study examines the contribution of CD8+CD28- cells to CTL memory pools for CTL clones specific for two persistent human viruses, CMV and HIV - clones were found to be similarly distributed in the CD28 depleted cell population.
- HIV CTL responses to 3 Env and 2 Gag peptides were studied.
- The clonal composition of the TCR  $V\beta$  responses was studied and was found to be highly focused, with one TCR beta-chain sequence tending to dominate the peptide-specific response clones to this epitope were  $V\beta$  13.6.

HXB2 Location gp160 (212-231)

Author Location gp120

Epitope PIPIHYCAPAGFAILKCNNK

Immunogen HIV-1 infection Species (MHC) human (B57)

References Jin et al. 1998b

- Progressive HIV-1 infection and CD4 decline was associated decreased the IL-2-expandable HIV-1 CTL response in 10 asymptomatic HIV-infected patients - this observation may be partially due to a reduction and impaired function of T helper cells, CTL exhaustion and APC dysfunction.
- Continued presence of HIV-1 specific memory cells (CTLp) was observed in three patients, one to GIWGCSGKLICTTAV, one to ALIWEDLRSLCLFSY, and one to PIPIHYCAPAG-FAILKCNNK.

HXB2 Location gp160 (212-231)

Author Location gp120

Epitope PIPIHYCAPAGFAILKCNNK

Immunogen HIV-1 infection

Species (MHC) human

References Weekes et al. 1999a

• Peptide 740.19: Memory CTL specific for HIV-1 may contribute to oligoclonal expansions within the CD57+ CD28- CD8+ CTLp populations.

HXB2 Location gp160 (212–231)

Author Location gp120 (219–238 HXB2)

Epitope PIPIHYCAPAGFAILKCNNK

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human

References Lieberman et al. 1992

• CTL epitope defined by T cell line and peptide mapping.

HXB2 Location gp160 (212-231)

**Author Location** gp120 (219–238)

Epitope PIPIHYCAPAGFAILKCNNK

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

• HIV-specific CTL lines developed by ex vivo stimulation with peptide.

HXB2 Location gp160 (217-226)

Author Location gp120 (217–226 HIV-MN)

Epitope YCAPAGFAIL

Epitope name YL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0102)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- This is a newly defined epitope. Last position (10)in the epitope had potentially experienced positive selection. YCAPAGFAIi escape variant was found.

HXB2 Location gp160 (237-246)

**Author Location** Env

**Epitope** GPCKNVSTVQ

**Immunogen** 

Species (MHC) human (B56)

References De Groot et al. 2001

- The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.
- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 predicted B7 superfamily (HLA B7, B8, and B58) epitopes were identified that could stimulate IFN $\gamma$  production in an ELISPOT assay.
- GPCKNVSTVQ was newly defined as an epitope in this study, was shown to stimulate an ELISPOT response, and to bind to HLA-B7.

HXB2 Location gp160 (239–247)

**Author Location** gp160 (237–245 BORI)

Epitope CKNVSTVQC

Epitope name gp160 CC9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (Cw\*0802)

**Donor MHC** A\*2902, B\*1402, Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

assav

Keywords dynamics, immunodominance, acute/early infection, kinetics, characterizing CD8+ T cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTLmediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient BORI had high viral loads and rapid CD4 decline. BORI mounted 8 detected responses. BORI did not control viral replication well, and escape mutations occurred early and in most epitopes, 6/7 that were precisely identified.
- Four variants of the CKNVSTVQC epitope were found in the patient BORI. CeNVSTVQC and cCeNVSTVhC came up first, at day 6 from onset of symptoms. The CeNVSTVQC form was the form that persisted, with a second rare variant present at day 35, CgNVSTVQC. These variants were not tested for their impact on escape.

HXB2 Location gp160 (239–247)

Author Location gp120 (241-249 LAI)

**Epitope** CTNVSTVQC

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw8)

References Sipsas et al. 1997

- HIV IIIB proteins were used to define the range of CTL epitopes recognized by 3 lab workers accidentally infected with HIV-1 IIIB.
- CTNVSTVQC contains a potential N-linked glycosylation site and cysteine residues, possibly related to a requirement for a high sensitizing dose of peptide for CTL activity.

HXB2 Location gp160 (242–261)

**Author Location** gp120 (249–268)

Epitope VSTVQCTHGIRPVVSTQLLL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

• HIV-specific CTL lines developed by ex vivo stimulation with peptide.

HXB2 Location gp160 (242-261)

Author Location gp120 (249–268 SF2)

Epitope VSTVQCTHGIRPVVSTQLLL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- · Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-2, -B21.

**HXB2 Location** gp160 (242–261) **Author Location** gp120 (249–268)

Epitope VSTVQCTHGIRPVVSTQLLL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

 CTL expanded ex vivo were later infused into HIV-1 infected patients.

**HXB2 Location** gp160 (252–260)

Author Location gp120 (255-263 SF2)

Epitope RPIVSTQLL Immunogen HIV-1 infection Species (MHC) human (B\*3501) References Tomiyama *et al.* 1997

- A CTL clone responsive to this epitope was obtained.
- Only 1/7 B35-positive individuals had a CTL response to this epitope.
- An I to V substitution at position 3 reduces specific lysis, but not binding to B\*3501.
- A Q to H substitution at position 7 abrogates specific lysis, but not binding to B\*3501.

**HXB2 Location** gp160 (252–260)

Author Location gp120 (255–263 SF2)

Epitope RPIVSTQLL Immunogen HIV-1 infection Species (MHC) human (B35) References Shiga *et al.* 1996

• Binds HLA-B\*3501.

**HXB2 Location** gp160 (252–260)

**Author Location (SF2)** 

Epitope RPIVSTQLL Immunogen HIV-1 infection Species (MHC) human (B35)

**Keywords** rate of progression **References** Kawana *et al.* 1999

- HLA B35 is associated with rapid disease progression.
- The sequences of 9 previously described HIV-1 B35 CTL epitopes were obtained in 10 HLA B35+ and 19 HLA B35- individuals.
- 3/9 CTL epitopes had substitutions that were more common in B35+ individuals than in B35- individuals, but this was one of the six that had no B35 associated pattern of mutation.

HXB2 Location gp160 (252–261)

Author Location Env

Epitope RPVVSTQLLL

Immunogen

Species (MHC) human (B7)

References De Groot et al. 2001

- The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.
- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 of the predicted B7 superfamily (HLA B7, B8, and B58) epitopes were identified that could stimulate IFNγ production in an ELISPOT assay.

 RPVVSTQLLL was one of the 15, and had been previously identified as an HLA-B7 epitope, and was confirmed in this study.

HXB2 Location gp160 (252–261)

Author Location Env

Epitope KPVVSTQLLL

**Epitope name** 1298 **Subtype** multiple

Immunogen HIV-1 infection Species (MHC) human (B7, B8)

**Donor MHC** A01, A03, B07, B08, Cw03, Cw07; A29, A30, B08, B44, Cw07, Cw16

Assay type T-cell Elispot

**Keywords** binding affinity, computational epitope prediction, cross-presentation by different HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for KPVVSTQLLL: 46% Promiscuous epitope binding to B08 and B07.

**HXB2 Location** gp160 (252–261)

**Author Location** Env

Epitope RPVVSTQLLL

Epitope name 1305
Subtype multiple

**Immunogen** HIV-1 infection **Species (MHC)** human (B7, B8)

Donor MHC A29, A30, B08, B44, Cw07, Cw16

Country United States. Assay type T-cell Elispot

**Keywords** binding affinity, supertype, computational epitope prediction, cross-presentation by differ-

ent HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for RPVVSTQLLL: 41%. Supertype epitope, published B07, responses by B08 subject.

**HXB2 Location** gp160 (252–271)

Author Location gp120 (256–275 LAI)

Epitope RPVVSTQLLLNGSLAEEEVV

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

References Shankar et al. 1996

HXB2 Location gp160 (252–271)

Author Location Env (256–268 BH10, LAI)

Epitope RPVVSTQLLLNGSLAEEEVV

Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is STQLLLNGSLAEE) has similarity with the lymphatic endothelium-specific hyaluronan receptor LYVE-1 fragment TTRLLVQGSLRAEE.

HXB2 Location gp160 (281–288)

**Author Location** (C consensus)

Epitope AKTIIVHL

Subtype C

**Immunogen** HIV-1 infection **Species (MHC)** human (Cw\*0602)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the L8 residue of AKTIIVHL are associated with the presence of the HLA presenting molecule in the host.
- AKTIIVHL not optimized.

**HXB2 Location** gp160 (291–307)

Author Location gp120 (295-312 BRU)

Epitope SVEINCTRPNNNTRKSI

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Dadaglio et al. 1991

• Defined through blocking CTL activity, and Env deletions.

**HXB2 Location** gp160 (291–307)

**Author Location** gp120 (291–307 IIIB)

Epitope SVEINCTRPNNNTRKRI

Subtype B

Immunogen vaccine

*Vector/Type:* DNA, DNA with protein boost *Strain:* B clade IIIB *HIV component:* gp160

Adjuvant: IL-12

Species (MHC) mouse (A2)

Keywords vaccine-specific epitope characteristics

References Kiszka et al. 2002

 Transgenic mice expressing a HLA-A2/Kb chimeric protein were vaccinated with a full length gp160 or with gp160deltaV3, with the V3 loop deleted. Mice given gp160deltaV3 had a broader immune response than those given gp160, with increased responses to conserved HLA-A2 epitopes in the C1 region of gp120, KLTPLCVTL, and the C-term region of gp41, SLLNATAIAV.

- Greater resistance was conferred by the gp160deltaV3 than the gp160 vaccine to a challenge of vaccinia expressing heterologous gp160 from primary isolates (VI-06 and 89.6), and the resistance was conferred by CD8+ T-cells.
- The most intense CTL responses to the intact gp160 vaccine were directed at three V3 peptides.

HXB2 Location gp160 (291–307)

Author Location Env (292-301 BH10, LAI)

Epitope SVEINCTRPNNNTRKSI

Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is VEINCTRPNN) has similarity with the FasI receptor precursor (Apptosismediating surface antigen fas) (APO-1 antigen) (CD95 antigen) fragment VEINCTRQN.

HXB2 Location gp160 (296–305)

Author Location Env

**Epitope** CTRPNNNTRK

Epitope name 1265

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3, A2)

**Donor MHC** A03, A23, B49, B57

Country United States.
Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion, cross-presentation by different HLA

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for CTRPNNNTRK: 51% Promiscuous epitope binding to A02 and A03.

 $\textbf{HXB2 Location} \hspace{0.1cm} gp160 \hspace{0.1cm} (297\text{--}322)$ 

Author Location gp120 (297–322 IIIB)

Epitope TRPNNNTRKRIRIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB HIV component: V3 Adjuvant: liposome

**Species (MHC)** mouse (H-2D<sup>d</sup>)

References Chang et al. 1999

- Induction of peptide-specific CTLs in BALB/c mice was dependent on immunization with peptide encapsulated liposomes containing MPL as adjuvant.
- T26K (26mer) elicited a stronger AB and CTL response than R15K (a V3 15mer, RIQRGPGRAFVTIGK)

**HXB2 Location** gp160 (297–330) **Author Location** Env (303–335 BX08)

Epitope TRPNNNTRKSIHIGPGRAFYATGEIIGDIRQAH

Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long amino acid peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I trial
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees – 6/10 reacted to this peptide.
- 9/12 tested mounted a CTL response to at least one of the six peptides; each of the six peptides elicited a CTL response in at least one individual – this peptide was particularly immunogenic, eliciting a CTL response in five vaccinees.
- None of the 12 tested had an IgG response to gp120 or gp160 and vaccinees could be differentiated from HIV-1 seropositive individuals with a commercial HIV detection kit – no neutralizing antibodies were observed.

**HXB2 Location** gp160 (298–307)

Author Location gp120 (298–307)

Epitope RPNNNTRKSI

Immunogen HIV-1 infection

Species (MHC) human (B\*07)

Keywords epitope processing, TCR usage

References Ferris et al. 1999; Hammond et al. 1995

- The processing of this epitope is TAP1/2-dependent, as are most Env epitopes, and it contains an N-linked glycosylation site that is glycosylated in Env.
- Peptide that had been deglycosylated, a process that changes asparagine (N) to aspartic acid (D) (RPNDNTRKSI) was recognized a 100-fold more efficiently than either glycosylated or non-glycosylated RPNNNTRKSI.
- Position 5 is not involved with HLA B\*07 binding, so is probably important for TCR recognition.
- HIV-1 Env epitopes are typically processed by a TAP1/2 dependent mechanism, which involves cotranslational translocation into the ER, glycosylation, export back into the cytosol, and deglycosylation for processing, and retransport into the ER for the association with class I molecules.
- The particular pathway of generating an epitope may have an impact on the presentation of that epitope, quantitatively as well as qualitatively.

HXB2 Location gp160 (298–307)

Author Location gp120 (302–312 HXB2)

Epitope RPNNNTRKSI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*0702 epitope.

**HXB2 Location** gp160 (298–307)

Author Location (C consensus)

Epitope RPNNNTRKSI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location gp160 (298-307)

Author Location gp120 (302–312 HXB2)

Epitope RPNNNTRKSI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Safrit et al. 1994b

• CTL from two acute seroconversion cases.

HXB2 Location gp160 (298–307)

**Author Location** gp120 (302–312 HXB2)

Epitope RPNNNTRKSI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Hammond et al. 1995

- Peptide processed by a TAP-1/2-dependent pathway only.
- CTL from an acute seroconverter.

HXB2 Location gp160 (298–307)

Author Location gp120 (302–312 HXB2)

Epitope RPNNNTRKSI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Wolinsky et al. 1996

• Longitudinal study of epitope variation in vivo.

**HXB2 Location** gp160 (298–307)

**Author Location** gp120 (302–311 subtype B)

Epitope RPNNNTRKSI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords subtype comparisons, immunodominance

References Wilson et al. 1998b

• The extent of CTL interclade cross-reactivity from CTL isolated from individuals newly infected with B clade virus was studied, and extensive cross-reactivity was observed.  Two HLA B7 individuals had CTL response to B\_LAI, A\_92UG037 and C\_92BR025 gp160, but were B clade strain MN non-responders – the authors note that the B7 epitope RPNNNTRKSI is immunodominant, conserved between the LAI and clade A and C strains, but is very divergent in MN (RPNYNKRKI), and that this epitope might be dominating the specificity of the response in the HLA B7 individuals.

HXB2 Location gp160 (298-307)

Author Location gp120 (303–312 SF2)

**Epitope** RPNNNTRKSI **Immunogen** HIV-1 infection **Species (MHC)** human (B7)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B7+ individuals that had a CTL response to this epitope broken down by group: 0/4 group 1, 1/3 group 2, and 1/1 group 3.

**HXB2 Location** gp160 (298–307)

Author Location gp120 (298–307) Epitope RPNNNTRKSI

Immunogen HIV-1 infection Species (MHC) human (B7)

**Keywords** rate of progression, acute/early infection **References** Day *et al.* 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

HXB2 Location gp160 (298–307) Author Location gp120 (298–307) Epitope RPNNNTRKSI Epitope name B7-RI10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 4/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 1/4 individuals had detectable responses to this epitope after STI.

HXB2 Location gp160 (298–307)

Author Location gp120

**Epitope** RPNNNTRKSI

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A2, A3, B7, Bw6

Keywords HAART, ART

References Appay et al. 2002

- Four HIV patients with prolonged clinically successful antiviral therapy but with ongoing evidence of replication and Nef mRNA transcription, showed specific T-cell responses by Elispot and Tetramer staining, maintained for 2-4 years after initiation of HAART.
- Nef epitope recognition was detected in all 4 subjects, gp120,
   Pol and Gag-specific in 1 or 2 subjects.
- The HIV-specific CD8+ T-cells had an intermediate maturation phenotype characterized by low levels of perforin and high levels of CD27 expression.

**HXB2 Location** gp160 (298–307)

Author Location gp160 (298–307)

Epitope RPNNNTRRGI

**Epitope name** B7-RI10 Env

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** supervised treatment interruptions (STI), escape, early treatment, superinfection

References Altfeld et al. 2002a

• An HIV+ individual given acute ARV and undergoing STI developed a strong CTL immune response, but despite this was shown to be superinfected with second B clade virus. Some epitope variants of the second infecting virus were resistant to the initial CTL response.

 The first infecting strain had the variant rpSnntrKSi, and the CTL response was higher to the second variant, RPNNNTR-RGI.

HXB2 Location gp160 (298-307)

Author Location gp120

**Epitope** RPNNNTRKSI **Immunogen** HIV-1 infection **Species (MHC)** human (B7)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** HIV exposed persistently seronegative (HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- No one pre-seroconversion, 0/9 HLA A2+ infection-resistant men, and 0/4 pre-seroconversion men who went on to become infected, reacted to this epitope.

**HXB2 Location** gp160 (298–307) **Author Location** Env (302–311)

Epitope RPNNNTRKSI Immunogen HIV-1 infection Species (MHC) human (B7)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 3/7 patients recognized this epitope.

**HXB2 Location** gp160 (298–307)

Author Location gp120 (303–312 IIIB)

Epitope RPNNNTRKSI Immunogen HIV-1 infection Species (MHC) human (B7?)

**Keywords** responses in children, mother-to-infant transmission

References Wilson et al. 1996

- Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.
- RPNNNTRKDI and RPNNNTRKGI, naturally occurring variants, were found in non-transmitting mother ability to recognize these variants has not yet been determined.

**HXB2 Location** gp160 (299–319)

**Author Location** Env (299–319)

Epitope PNNNTRKSIRIGPGQTFYA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons
References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location gp160 (303–322)

Author Location gp120

Epitope TRKSIHIGPGRAFYTTGE

Immunogen vaccine

Vector/Type: virus-like particle (VLP)
Strain: B clade consensus HIV component:
Gag, V3

Species (MHC) mouse

References Luo et al. 1998

 Intramuscular injection of chimeric gag-env virus-like particles (VLPs) containing V3 loop sequences into BALB/c mice induce V3 specific CTL – TRKSIHIGPGRAFYTTGE is a B subtype consensus that stimulated a cross-reactive CTL response.

**HXB2 Location** gp160 (304–318)

Author Location gp120 (304-318 IIIB)

Epitope RKSIRIQRGPGRAFV

Immunogen vaccine

Vector/Type: virus-like particle (VLP) Strain: B clade IIIB, B clade MN, B clade RF, B clade SF2, HIV-2 VLP HIV compo-

nent: Gag, V3

**Species (MHC)** mouse (H-2<sup>d</sup>) **References** Kang *et al.* 1999

- Virus-like particles could be formed from HIV-2 gag after deleting 143 amino acids at the C-terminal end a proline rich region in positions 373-377 was critical to VLP formation.
- CTL responses in BALB/c mice were induced by chimeric gag-V3 particles against the V3 region of HIV-1 clade B isolates IIIB (SIRIQRGRAFVTI), MN (KRIHIGPGRAFYTTK), RF (SITKGPGRVIYATGQ), and SF2 (SIYIGPGRAFHTTGR)
- The vaccine induced CTL were cross-reactive with a broad spectrum of B clade isolates, with the exception of the RF V3 which did not induce CTL.

 $\textbf{HXB2 Location} \hspace{0.1cm} \texttt{gp160} \hspace{0.1cm} (305\text{--}321)$ 

Author Location gp120 (MN)

Epitope KRIHIGPGRAFYTTK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A2

Assay type CD8 T-cell Elispot - IFNγ

References Chitnis et al. 2003

• 17 perinatally HIV-1 infected children (0.08-16 years) were evaluated for HLA-A2-restricted IFN-gamma CD8+ CTL responses against 4 immunodominant peptides that carry HLA-A2 epitopes. Two peptides were from gp120 (one at position 112, one from the V3 loop), and one each was from gp41 and Gag. 15/17 patients responded to the Gag peptide, 13/17 to the gp41 and the non-V3 gp120 peptides, and 11/17 responded to the V3 loop. 4 children recognized all 4 peptides.

HXB2 Location gp160 (306-322) Author Location gp160 (LAI)

Epitope SIRIQGPGRAFVTIGI

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI HIV component: gp160 Adjuvant: aluminum hydroxide, CpG immunostimulatory sequence (ISS)

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Keywords immunodominance, Th1, Th2

References Deml et al. 1999

• Addition of CpG oligodeoxynucleotide to a gp160/alum vaccine given to BALB/c mice shifted the response to Th0/Th1 from Th2, but no still CTL response to this immunodominant epitope was induced.

HXB2 Location gp160 (307-324)

**Author Location** (C consensus)

Epitope IRIGPGQTFYATGDII

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression References Kiepiela et al. 2007

• A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location gp160 (308-321)

**Author Location** Env (IIIB)

Epitope RIQRGPGRAFVTIG

Epitope name P18IIIB

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

HIV component: V3

Species (MHC) mouse (Dd)

Keywords binding affinity, Th1

References Ahlers et al. 2001

• BALB/c and A.AL were immunized with an Env-peptide vaccine construct containing the CTL epitope P18IIIB and the T helper epitope T1, KQIINMWQEVGKAMYA.

- **Keywords** assay standardization/improvement, HAART, Substitution of Glu (wt) to Ala in T1, kqiinmwqAvgkamya, caused increased affinity for MHC class II Ek, resulting in the upregulation of CD40L in the responding Th cells, and shifting the response towards Th1. Increased Th responses stimulated DCs to produce higher levels of IL-12, and B7-1 and B7-2, and enhanced CTL responses to P18.
  - The modified epitope, T1A, elicited stronger protection against increasing doses of viral challenge with vaccinia expressing HIV-1 IIIB gp120 compared to the wt epitope T1.

HXB2 Location gp160 (308–321)

**Author Location** Env (gp160)

Epitope RIQRGPGRAFVTIK

Epitope name P18IIIB Immunogen vaccine

Vector/Type: hemagglutinating virus of Japan

(HVJ)-liposome Strain: B clade IIIB HIV

component: gp160

Species (MHC) mouse Donor MHC H-2d

Assay type Cytokine production, Chromium-release as-

sav

Keywords genital and mucosal immunity

References Sakaue et al. 2003

- BALB/c mice were immunized nasally with HIVgp160encapsulated hemagglutininating virus of Japan (HVJ)liposome. Vaccination induced IgG in serum and IgA in nasal wash, saliva, fecal extract, and vaginal wash, with some ability to neutralize the primary field isolate HIV-MNp.
- Th1 and Th2-type responses were stimulated, as well as gp160 V3-specific MHC class I-restricted CTL responses.

HXB2 Location gp160 (308–322)

**Author Location** Env (315–329)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Subtype B

Immunogen vaccine

Vector/Type: DNA HIV component: HIV-1

Species (MHC) mouse (A\*0201)

Keywords epitope processing, vaccine-specific epitope

characteristics, immunodominance

References Singh et al. 2002; Sykes & Johnston 1999

- C3H (H-2k) transgenic mice carrying a fused HLA-A\*0201 alpha1 and alpha2 and H-2Dk alpha3 hybrid class I molecule were immunized using an epidermal gene gun with an ubiquitin expression library of 32 plasmids that spanned the HIV-1 genome. Ubiquitin targets the expressed HIV-1 peptides to the proteasome.
- A single immunization with the UB-HIV-1 library vaccine induced potent, stable and multivalent CTL responses against all library members.
- Immunodominant epitopes SLYNTVATL (Gag), ILKEPVHGV (Pol), RIQRGPGRAFVTIGK (Env) and AFHHVAREK (Nef) elicited strong CD8+/IFN- responses and stimulated CTL that were functional in a Cr-release assay and against wild type antigen.

 The presence of multiple plasmids HLA-A\*0201-restricted CTL epitopes did not decrease CTL immunogenicity, and CTL responses to single peptide immunizations were comparable to responses based on mixtures of either 16 or 32 peptides.

HXB2 Location gp160 (308–322) Author Location gp120 (315–329 IIIB) Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species (MHC)** human (A11) **References** Achour *et al.* 1994

• One of 3 HLA type restrictions associated with this peptide.

HXB2 Location gp160 (308–322)
Author Location gp120 (315–329 BRU)
Epitope RIQRGPGRAFVTIGK
Immunogen HIV-1 infection
Species (MHC) human (A2)
References Dadaglio et al. 1991

• Defined through blocking CTL activity, and Env deletions.

HXB2 Location gp160 (308–322)
Author Location gp120 (315–329 IIIB)
Epitope RIQRGPGRAFVTIGK
Immunogen HIV-1 infection
Species (MHC) human (A2)
References Clerici et al. 1991a

 Helper and cytotoxic T cells can be stimulated by this peptide (P18)

HXB2 Location gp160 (308–322) Author Location gp120 (308–322 IIIB) Epitope RIQRGPGRAFVTIGK

**Subtype** B **Immunogen** vaccine

Vector/Type: DNA, DNA with protein boost Strain: B clade IIIB HIV component: gp160

Adjuvant: IL-12

Species (MHC) mouse (A2)

Keywords vaccine-specific epitope characteristics

References Kiszka et al. 2002

- Transgenic mice expressing a HLA-A2/Kb chimeric protein were vaccinated with a full length gp160 or with gp160deltaV3, with the V3 loop deleted. Mice given gp160deltaV3 had a broader immune response than those given gp160, with increased responses to conserved HLA-A2 epitopes in the C1 region of gp120, KLTPLCVTL, and the C-term region of gp41, SLLNATAIAV.
- Greater resistance was conferred by the gp160deltaV3 than the gp160 vaccine to a challenge of vaccinia expressing heterologous gp160 from primary isolates (VI-06 and 89.6), and the resistance was conferred by CD8+ T-cells.
- The most intense CTL responses to the intact gp160 vaccine were directed at three V3 peptides.

HXB2 Location gp160 (308–322) Author Location gp120 (315–329 IIIB) Epitope RIQRGPGRAFVTIGK Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (A2, A3)

References Achour et al. 1993

• Two of 3 HLA type restrictions associated with this peptide.

HXB2 Location gp160 (308–322) Author Location gp160 (308–322) Epitope RIQRGPGRAFVTIGK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

**HXB2 Location** gp160 (308–322) **Author Location** gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB HIV component: V3

Species (MHC) mouse (Dd)

References Takahashi et al. 1989a

Positions R(8) and F(10) are important for MHC/peptide interaction.

HXB2 Location gp160 (308–322) Author Location gp120 (315–329 IIIB) Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

HIV component: V3

Species (MHC) mouse (D<sup>d</sup>)

References Sastry et al. 1992

 Free peptide injected into the footpad of a mouse could stimulate specific CTL.

HXB2 Location gp160 (308–322) Author Location gp120 (315–329 IIIB) Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN

HIV component: V3

**Species (MHC)** mouse (D<sup>d</sup>)

References Ahlers et al. 1997b

- PCLUS 3-18MN synthetic peptide vaccine construct contained T1 helper epitope covalently linked to truncated P18 CTL epitope.
- A substitution in the T1 peptide stimulated an enhanced Th response and class II binding specificity, which in turn enhanced CTL induction by vaccine.
- Construct PCLUS 3-18MN is currently in a phase I vaccine clinical trial.

HXB2 Location gp160 (308–322) Author Location gp120 (313–327 MN) Epitope RIHIGPGRAFYTTKN

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB, B clade MN HIV component: gp160

**Species (MHC)** mouse (D<sup>d</sup>)

References Takahashi et al. 1989b

• Y(11 MN) exchange with V(11 IIIB) interchanges specificities.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (313–327 IIIB, MN, RF)

Epitope SITKGPGRVIYATGQ

Immunogen vaccine

HIV component: gp160

**Species (MHC)** mouse (D<sup>d</sup>)

References Takahashi et al. 1992

• Comparison of MN, IIIB, and RF specificities, position 11 is critical.

HXB2 Location gp160 (308-322) Author Location gp160 (315–329 IIIB) Epitope RIQRGPGRAFVTIGK

**Epitope name** P18

Immunogen in vitro stimulation or selection

Species (MHC) mouse (Dd) Donor MHC H-2d Keywords TCR usage

References Yokosuka et al. 2002

· The TCR repertoire and its specificity was studied through analyzing the spectrum of TCR-alpha and beta chains able to reconstitute a reaction to the H-2 Dd-restricted P18 peptide. The RT-1 TCR alpha chain was able to react with 1/3 of the tested TCR beta chains to create a specific response. Experiments in transgenic mice also supported the observation that a single TCR alpha chain would confer the specificity of the response and could interact with a large variety of TCR beta

HXB2 Location gp160 (308-322) **Author Location** gp120 (HXB2) **Epitope** RIQRGPGRAFVTIGK

> Subtype B Immunogen vaccine

> > Vector/Type: protein HIV component: Gag,

**Species (MHC)** mouse (H-2<sup>d</sup>) References Griffiths et al. 1993 • Gag-V3 fusion protein immunization elicited V3 CTL response in mice.

HXB2 Location gp160 (308-322) Author Location gp120 (HXB2) Epitope RIQRGPGRAFVTIGK

Subtype B

Immunogen vaccine

Vector/Type: virus-like particle (VLP) HIV

component: Env, Gag

Species (MHC) mouse (H-2<sup>d</sup>)

References Deml et al. 1997

• Env bound to virus-like particles (VLPs) can elicit a CTL response that is dependent on the amount of Env presented on the VLP.

**HXB2 Location** gp160 (308–322) Author Location gp120 (313–327 MN) Epitope RIHIGPGRAFYTTKN

Immunogen vaccine

Vector/Type: DNA Strain: B clade MN HIV component: gp160, V3

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Fomsgaard et al. 1998a

Vector/Type: vaccinia Strain: B clade RF • Enhanced B and CTL responses to the V3 region occur following epidermal immunization by gene gun with a chimeric DNA vaccine of V3-hepatitis B surface antigen relative to a gp160 plasmid vaccine.

> HXB2 Location gp160 (308–322) Author Location gp120 (313-327 MN) Epitope RIHIGPGRAFYTTKN

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN HIV component: V3 Adjuvant: GM-CSF,

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1

References Ahlers et al. 1996; Ahlers et al. 1997a

- · Vaccine constructs containing helper, antibody and CTL peptide epitopes induce strong Th1, CTL and NAb responses against the autologous HIV-1 virus.
- The peptide CTL response was as cross-reactive as one elicited by a vaccinia construct expressing rgp160 MN.
- GM-CSF and IL-12 were the two cytokines most effective for inducing and boosting CTLs.

HXB2 Location gp160 (308–322) Author Location gp120 (315–329 IIIB) Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

*Vector/Type:* virus-like particle (VLP) Strain: B clade IIIB HIV component: Gag, V3

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Layton et al. 1993

• V3-Ty-Virus-like particles can induce type-specific CTL in mice in the absence of adjuvant.

HXB2 Location gp160 (308-322)

Author Location gp120 (IIIB)

Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB HIV component: gp120 Adjuvant: IL-2, IL-

2/Ig

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Barouch et al. 1998

- A discistronic IL-2 gp120 expression vector gave a weaker CTL response than gp120 alone in the expression vector, however co-administration of an II-2/IgG fusion protein enhanced the immune response and administration of a II-2/IgG plasmid had a response that depended on the timing of administration.
- This study showed that a response to an HIV-1 DNA vaccine could be either augmented or suppressed by plasmid Cytokine/Ig administration.

**HXB2 Location** gp160 (308–322)

**Author Location** Env (308–322 IIIB) **Epitope** RIQRGPGRAFVTIGK

Epitope name P18
Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB HIV component: V3 Adjuvant: B7, CpG immunostimulatory sequence (ISS), in vivo electroporation

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1

References Uno-Furuta et al. 2001

- Peptide immunization usually doesn't elicit a good CTL response because epitopes are not internalized and processed and presented, so vaccination with electric pulsing was tried (i.m. injection followed by 8 electric pulses), to enhance peptide uptake through electroporation.
- BALB/c immunized with HIV P18 or hepatitis C P17 peptides with an electric pulse elicited a CTL response, those that did not receive the pulse did not.
- The CTL response was enhanced by addition of immunostimulatory sequences ISS in the plasmid pCMV-LacZ, that contains hexamers GACGTC, AGCGCT, AACGCT, sequences common in prokaryotic genomes but rare in eukaryotic genomes that elicit Th1 cytokines and result in B cell and T-cell proliferation.
- The CTL response was also enhanced by addition of B7-1 cDNA – the B7 family of proteins transduce co-stimulatory signals through interaction with CD28.

**HXB2 Location** gp160 (308–322)

**Author Location** gp160 (MN)

 ${\bf Epitope} \ \ {\tt RIHIGPGRAFYTTKN}$ 

Immunogen vaccine

Vector/Type: DNA Strain: B clade MN

HIV component: gp160

**Species (MHC)** mouse  $(H-2^d, H-2^b)$ 

References Fomsgaard et al. 1998b

 CTL responses to a primary gene gun vaccination were rapid and strong for several methods of vaccinations: i.m., bupivacaine pretreatment, cardiotoxin pretreatment or gene gun – the CTL response was more rapid and consistent than the antibody response. **HXB2 Location** gp160 (308–322) **Author Location** gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>p</sup>, H-2<sup>u</sup>, H-2<sup>q</sup>)

References Shirai et al. 1992; Shirai et al. 1993

- In a murine system multiple class I molecules can present this
  peptide, called P18, to CTL, including H-2D<sup>d</sup>, H-2D<sup>p</sup>, H-2D<sup>q</sup>,
  H-2L<sup>q</sup>
- The MHC class I molecule D<sup>d</sup> as well as H-2<sup>u,p,q</sup>, were found to present peptides P18 and HP53.
- The V-β usage in T cells showing cross-reaction between these two peptides was conserved for H-2<sup>d,u,p</sup>, but not in H-2<sup>q</sup>

HXB2 Location gp160 (308–322)

Author Location gp160 (IIIB)

Epitope GIHIGPGRAFYAARK

Immunogen vaccine

Vector/Type: peptide, protein Strain: B clade IIIB HIV component: gp160 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Keywords Th1, Th2

References Morris et al. 2000

 LT(R192G) induces gp160-specific serum and mucosal IgG1 and IgG2a, systemic CTL activity and Th1 and Th2 cytokine responses upon internasal immunization.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (315-329 IIIB)

Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB HIV component: V3 Adjuvant: Cholera tovin (CT)

toxin (CT)

Species (MHC) mouse (H-2D<sup>d</sup>)

References Porgador et al. 1997

- A intranasal peptide vaccine with cholera toxin as a mucosal adjuvant was given.
- IIIB peptide referred to as R15K.
- Peptide-specific CTLs were induced after *in vitro* restimulation with peptide-pulsed targets.
- R15K was superior at inducing CTL compared to the RGP-GRAFVTI, in contrast to the findings of Nehete *et al*.
- Memory CTL responses were induced.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (315-329 IIIB)

Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: vaccinia with H1 influenza HA gene cassette Strain: B clade IIIB HIV

component: p18 Gag

Species (MHC) (H-2D<sup>d</sup>)

References Chiba et al. 1999

BALB/c mice, but could not induce a P18IIIB-specific antibody response.

HXB2 Location gp160 (308-322) Author Location gp120 (multiple) Epitope RIHIGPGRAFYTTKN

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN,

B clade SC HIV component: V3

**Species (MHC)** mouse (H-2D<sup>d</sup>) References Casement et al. 1995

• V3 peptides from MN and SC induce murine CTL that are cross-reactive with diverse strains.

**HXB2 Location** gp160 (308–322) Author Location gp120 (313–327 MN) Epitope RIHIGPGRAFYTTKN

Immunogen vaccine

Vector/Type: protein Strain: B clade MN HIV component: gp120 Adjuvant: QS21

**Species (MHC)** mouse (H-2D<sup>d</sup>) References Newman et al. 1997

· MN vaccine induced CTL reactive with MN, IIIB and RF vaccinia-expressed Env, but not this peptide.

HXB2 Location gp160 (308-322) Author Location gp120 (315–329 IIIB) Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp120 Adjuvant: QS21

**Species (MHC)** mouse (H-2D<sup>d</sup>) References Newman et al. 1997

• IIIB vaccine induced IIIB type-specific CTL to this peptide (P18), and an additional Env CTL response that was crossreactive.

**HXB2 Location** gp160 (308–322) Author Location gp120 (315–329) **Epitope** RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2D<sup>d</sup>) References Takahashi et al. 1988

V3 loop CTL response in mice vaccinated with gp160.

HXB2 Location gp160 (308-322) Author Location gp120 (315-329) Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: liposome Strain: B clade IIIB HIV component: V3 Adjuvant: oligomannose

**Species (MHC)** mouse (H-2D<sup>d</sup>)

References Fukasawa et al. 1998

• The peptide RIQRGPGRAFVTIGK was incorporated into liposomes and given as a subcutaneous injection, which induces a MHC class I restricted CTL response in mice.

• Vaccine was capable of priming P18IIIB specific CTL in • Liposomes coated with oligomannose show no toxicity and can elicit a potent CTL response upon a single subcutaneous infection, while non-coated liposomes do not, suggesting that oligomannose may be a good adjuvant for CTL responses.

HXB2 Location gp160 (308–322)

**Author Location** 

Epitope RIQRGPGRAFVTIGK

Epitope name P18 Subtype B Immunogen vaccine

> Vector/Type: fusion protein with anthrax delivery domain HIV component: V3 Adjuvant: B. anthracia lethal toxin LF component

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Keywords epitope processing, vaccine-specific epitope characteristics

References Lu et al. 2000a

· Bacillus anthrax lethal toxin (LFn)-HIV fusion proteins are candidate HIV vaccines that are safe in mice, and LFn-V3 region fusion proteins induce CD8 T cells in BALBc mice. LFn causes exogenous protein to be taken up and processed in a class I pathway. Expressed proteins from Gag p24 and nef fragments cloned into the LFn expression plasmid stimulate gag-specific CD4 proliferation and CTL responses in HIVinfected donor PBMCs in vitro.

HXB2 Location gp160 (308–322) Author Location gp120 (V3) (MN) **Epitope RIHIGPGRAFYTTKN** 

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB HIV component: V3 Adjuvant: Cholera toxin (CT), GM-CSF, IL-12, IL-18, IL-1 $\alpha$ 

**Species (MHC)** mouse (H-2D<sup>d</sup>) References Staats et al. 2001

- Cholera toxin (CT) is a potent adjuvant used in animal studies that is not safe in humans, so combinations of cytokins were used in nasal immunization of BALB/c mice V3 peptides to attempt to replace CT as a potent adjuvant.
- · Peptide vaccine induced CTL activity was significantly increased by IL-1alpha, IL-18, and GMCSF given alone as adjuvant, but CT gave more potent CTL activity than any single
- Combinations of cytokins could be more potent that CT as an adjuvant. The highest tetramer binding of H-2Dd peptidespecific PBMC after nasal immunization was observed with IL-1alpha plus IL-18 as adjuvant.
- Nasal immunization with HIV peptide in the presence of IL-1alpha, IL-12 and GM- CSF induced IFN-gamma-secreting cells in the cervical lymph node, the lung and the spleen, and was associated with upregulation of MHC class II and B7.1 on nonlymphocytes in NALT/nasal mucosal cells.
- Consistent results were obtained for the IIIB and the MN peptides.

HXB2 Location gp160 (308–322) Author Location gp160 (315–329 MN) Epitope RIHIGPGRAFYTTKN

Epitope name P18

Immunogen in vitro stimulation or selection

**Species (MHC)** mouse (H-2D<sup>d</sup>)

**Donor MHC** H-2d **Keywords** TCR usage

References Yokosuka et al. 2002

• The TCR repertoire and its specificity was studied through analyzing the spectrum of TCR-alpha and beta chains able to reconstitute a reaction to the H-2 Dd-restricted P18 peptide. The RT-1 TCR alpha chain was able to react with 1/3 of the tested TCR beta chains to create a specific response. Experiments in transgenic mice also supported the observation that a single TCR alpha chain would confer the specificity of the response and could interact with a large variety of TCR beta chains.

**HXB2 Location** gp160 (308–322)

Author Location Env (IIIB)

Epitope RIQRGPGRAFVTIGK

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp120 Adjuvant: poly(I:C), lipopolysaccharide (LPS)

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Assay type Chromium-release assay

Keywords epitope processing, vaccine-induced epitopes,

Th1, Th2, immunotherapy, adjuvant compari-

son

References Fujimoto et al. 2004

When BALB/c mice were immunized with recombinant HIV-1
 Env gp120 or Influenza HA protein together with polyriboinosinic polyribocytidylic acid (poly (I:C)), an epitope-specific CD8+ class I MHC-restricted CTL response was observed. This response was not observed when LPS was used as adjuvant instead of poly (I:C) indicating activation of cellular immunity by poly (I:C). In the presence of poly (I:C), immature DC presented processed external antigen in association with class I MHC.

**HXB2 Location** gp160 (308–322)

**Author Location** gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

**Species (MHC)** mouse (H-2D<sup>d</sup>, H-2D<sup>p</sup>, H-2D<sup>q</sup>, H-2<sup>u</sup>)

References Shirai et al. 1996b

 Multiple murine MHC can cross-present this epitope (P18) and HP53, DRVIEVVQGAYRAIR, to specific CTL.

**HXB2 Location** gp160 (308–322)

Author Location gp160 (MN)

Epitope RIHIGPGRAFYTTKN

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN HIV component: V3 Adjuvant: Montanide

(ISA 51)

Species (MHC) human

References Pinto et al. 1999

- Peptide P18: Eight HIV+ individuals were vaccinated with peptides containing specific T helper, CTL and Ab epitopes in Montanide ISA 51 in a Phase I trial.
- Four displayed a 4-fold increase in PCLUS 3-18 MN-specific T helper responses.
- One patient developed a new, sustained P18MN-peptide-specific CTL response the patient's HLA haplotype was A2,30; B53,7; Cw2,4, and anti-HLA A2 antibody did not inhibit the response, suggesting it was not A2.
- Patients with low baseline Ab levels developed an increase of neutralizing Ab titers.
- No significant change was observed in plasma HIV viral loads and CD4 cell counts.

HXB2 Location gp160 (308-322)

Author Location gp120 (MN)

**Epitope** RIHIGPGRAFYTTKN

Immunogen HIV-1 infection

Species (MHC) chimpanzee

References Lubeck et al. 1997

- Epitope-specific CTL detected in chimpanzees immunized with adenovirus-HIV-1 MN gp160 recombinant.
- CTL response may account for protection against subsequent HIV-1 SF2 challenge in a chimpanzee lacking neutralizing antibodies.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Immunogen HIV-1 exposed seronegative

Species (MHC) human

References Pinto et al. 1995

 CTL and T helper cell reactivity in healthcare workers exposed to HIV.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (313-327 MN)

Epitope RIHIGPGRAFYTTKN

Immunogen HIV-1 exposed seronegative

Species (MHC) human

References Pinto et al. 1995

 CTL and T helper cell reactivity in healthcare workers exposed to HIV.

**HXB2 Location** gp160 (308–322)

**Author Location** gp120 (110–122)

Epitope RIQRGPGRAFVTIGK

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB Adjuvant: FLt3 ligand (FL), GM-CSF, IL-12,

IL-15, IL-2

Species (MHC) mouse

Keywords vaccine-specific epitope characteristics

References Moore et al. 2002a

Intramuscular immunization of BALB/c mice with DNA vaccines carrying either gp160 or Nef in the expression vector plasmid pNGVL gave different responses – gp160 induced strong gp160-specific CTL and IFN-responses and low-titer humoral responses, and Nef generated humoral (IgG1, IgG2a) responses and IFN-responses but little CTL activity.

- Co-injection of DNA plasmids encoding cytokines and/or hematopoietic growth factors, IL2, IL-12, IL-15, Flt3 ligand (FL), and GMCSF tended to give responses that were enhanced quantitatively, but not altered qualitatively.
- · Co-administration of GMCSF most strongly enhanced CTL and IFN-responses against pNGVL-gp160.
- Repeated immunization with pNGVL-Nef failed to induce CTL responses. Co-administration of IL-12 most strongly enhanced humoral and IFNgamma responses.
- FL, which enhances innate immune responses, in combination with IL-2, IL-12 or IL-15 generated with most potent Nef responses.

HXB2 Location gp160 (308-322)

Author Location gp140 (iiib)

Epitope RIQRGPGRAFTIGK

Subtype B

Immunogen vaccine

Vector/Type: liposome, protein Strain: B clade IIIB HIV component: oligomeric gp140 Adjuvant: liposome

Species (MHC) mouse

Donor MHC H-2d

Assay type proliferation, Chromium-release assay

Keywords adjuvant comparison References Richards et al. 2004

• Mice were immunized with gp140 and an adjuvant that was an oil-in -water emulsion containing liposomes with lipid A with encapsulated antigen. Stable and unstable emulsions were found to have similar potencies of inducing antigen-specific T-cell proliferation and IgG antibodies, but stable emulsions also induced antigen-specific CTL responses. Stable emulsions had lowered IgG2a/IgG1 ratios than unstable.

HXB2 Location gp160 (309–317)

Author Location gp120 (310–318 SF2)

Epitope IYIGPGRAF Immunogen HIV-1 infection Species (MHC) human (A\*2402)

References Ikeda-Moore et al. 1997

- Defined using reverse immunogenetics 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) – 53 of the 59 peptides bound A\*2402.
- This peptide induced CTL in 1/4 HIV-1 + people tested.
- IYIGPGRAF bound to A\*2402 strongly, the epitope can be processed in a vaccinia construct and presented – no specific CTL clones were obtained.

**HXB2 Location** gp160 (309–318)

Author Location gp120 (314–323 CM243 subtype CRF01)

Epitope ITVGPGQVFY

Epitope name E309-318

Subtype CRF01\_AE Immunogen HIV-1 infection

Species (MHC) human (A11)

Keywords HIV exposed persistently seronegative

(HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers - weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not
- This epitope was strongly reactive in HIV+ control study subject 184 who carried HLA-A11.

**HXB2 Location** gp160 (309–318)

**Author Location** gp120 (314–323 CM243 subtype CRF01)

**Epitope** ITVGPGQVFY Subtype CRF01\_AE Immunogen HIV-1 infection Species (MHC) human (A11)

Keywords subtype comparisons

References Bond et al. 2001

- HLA-A11 CRF01 (called subtype E in Bond et al.) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This is one of the new A11 epitopes identified through the streamlined EpiMatrix method, and 2/8 tested FSWs recog-
- This epitope was not conserved in other subtypes, and exact matches were rare.

HXB2 Location gp160 (310-318)

**Author Location** 

Epitope HIGPGRAFY

**Epitope name** Env-HY9

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (A\*3002)

Donor MHC A\*3002 A\*3201 B\*4501 B\*5301 Cw\*0401 Cw\*1202

Kevwords HAART, ART References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- This epitope was newly defined in this study.
- · Subject 00RCH33 was on HAART had a viral load of 2900 and CD4 count of 727 and also recognized the epitopes YPLTFG-WCY, Nef(135-143), HLA B\*5301; AETFYVDGA, RT(437-445), HLA B\*4501; and RSLYNTVATLY, p17(76-86), HLA
- Among HIV+ individuals who carried HLA A30, 3/16 (19%) recognized this epitope.

HXB2 Location gp160 (310–318)
Author Location gp120 (310–318)
Epitope HIGPGRAFY
Immunogen HIV-1 infection
Species (MHC) human (A\*3002)
Keywords optimal epitope
References Frahm et al. 2007

HXB2 Location gp160 (310-318)

**Author Location** 

**Epitope** HIGPGRAFY **Epitope name** Env-HY9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2) References Sabbaj *et al.* 2003

Among HIV+ individuals who carried HLA A02, 6/29 (21%) recognized this epitope.

HXB2 Location gp160 (310–318)

Author Location gp160 (313–321 WEAU)

**Epitope** TLGPGRVLY **Epitope name** gp160 TY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*2902, B\*4403, B\*0801

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, immunodominance, acute/early infection, kinetics, characterizing CD8+ T cells, reversion, viral fitness

reversion, virai nu

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance. WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.
- This was one of five reasonably strong responses in early infection in the patient WEAU, and the epitope sequence did not vary during the first year of the infection.

**HXB2 Location** gp160 (310–323)

Author Location gp120 (315–328 MN)

Epitope HIGPGRAFYTTKNI

Epitope name p97
Immunogen vaccine

*Vector/Type:* canarypox prime with pseudovirion boost *Strain:* B clade IIIB, B clade MN *HIV component:* Gag, gp120, Protease

**Species** (MHC) mouse (H-2D<sup>d</sup>) **References** Arp *et al.* 1999

 The vaccine vCP205, canarypox vector, MN gp120 + Gag/Pro IIIB, with a HIV-1 pseudovirion boost was given to mice;)

• HIV-1 pseudovirion boost enhanced the CTL to this epitope in immunized BALB/c mice as measured by CTL lysis and IFN gamma production.

HXB2 Location gp160 (311–318)

**Author Location (MN)** 

Epitope IGPGRAFY

Immunogen vaccine

Vector/Type: B. abortus complex Strain: B clade MN HIV component: V3

**Species (MHC)** mouse (H-2D<sup>d</sup>)

References Golding et al. 2002a

Intranasal immunization of B. abortus conjugated to V3 peptides induces mucosal IFN-gamma producing T-cell responses in BALB/c mice.

**HXB2 Location** gp160 (311–319)

Author Location gp120 (311-320 IIIB)

Epitope RGPGRAFVT

Subtype B

Immunogen vaccine

Vector/Type: DNA, DNA with protein boost Strain: B clade IIIB HIV component: gp160

Adjuvant: IL-12

Species (MHC) mouse (A2)

**Keywords** vaccine-specific epitope characteristics

References Kiszka et al. 2002

- Transgenic mice expressing a HLA-A2/Kb chimeric protein were vaccinated with a full length gp160 or with gp160deltaV3, with the V3 loop deleted. Mice given gp160deltaV3 had a broader immune response than those given gp160, with increased responses to conserved HLA-A2 epitopes in the C1 region of gp120, KLTPLCVTL, and the C-term region of gp41, SLLNATAIAV.
- Greater resistance was conferred by the gp160deltaV3 than the gp160 vaccine to a challenge of vaccinia expressing heterologous gp160 from primary isolates (VI-06 and 89.6), and the resistance was conferred by CD8+ T-cells.
- The most intense CTL responses to the intact gp160 vaccine were directed at three V3 peptides.

**HXB2 Location** gp160 (311–319)

Author Location gp120 (312–320 SF2)

Epitope IGPGRAFHT

Immunogen vaccine

Vector/Type: DNA Strain: B clade SF2 HIV component: gp120

**Species (MHC)** mouse (D<sup>d</sup>)

References Selby et al. 1997

 Murine CTL response to peptide observed after immunization with DNA plasmid containing HIV-1 (SF2) gp120 gene regulated by bacteriophage T7 promoter.  CTL response required coadministration of rec vaccinia virus expressing T7 RNA polymerase or T7 RNA polymerase soluble protein.

HXB2 Location gp160 (311–319) Author Location gp120 (SF2) Epitope IGPGRAFHT Immunogen vaccine

*Vector/Type:* DNA prime with gp120 boost *Strain:* B clade SF2 *HIV component:* gp120

**Species (MHC)** mouse (H-2D<sup>d</sup>) **References** Barnett *et al.* 1997

- CTL were induced by vaccine, and restimulated *in vitro* with V3 peptide.
- DNA vaccine with protein boost stimulated both CTL and antibodies.
- Strains SF2 (IGPGRAFHT), US4 (IGPGRAFYA), and CM235 (IGPGQVFYR) were tested.

**HXB2 Location** gp160 (311–319) **Author Location** gp120 (312–320 SF2)

Epitope IGPGRAFHT

**Subtype** B **Immunogen** vaccine

Vector/Type: DNA, vaccinia Strain: B clade

SF2 HIV component: Gag, gp120 Species (MHC) mouse (H-2D<sup>d</sup>)

Assay type Chromium-release assay

Keywords epitope processing, vaccine-induced epitopes

References Doe et al. 1996

 Spleen cells from mice with distinct MHC types were infused into HIV vaccinated scid mice, to study the antigen presenting cells used by CTL induced in intramuscular injections. Bone marrow derived cells are used for presenation, but DNA infection is not required for priming, rather APCs can present proteins synthesized in other host cells.

**HXB2 Location** gp160 (311–320) **Author Location** gp160 (318–327 IIIB)

Epitope RGPGRAFVTI

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

References Alexander-Miller et al. 1996

- This epitope stimulates a CTL line derived from an HIV negative donor.
- This immunogenic peptide does not have the known binding motif for A2.1.
- The same optimal peptide for this human HLA-A2.1 epitope was observed for a murine H-2 D<sup>d</sup> epitope.

**HXB2 Location** gp160 (311–320)

**Author Location** gp120 (311–320 IIIB) **Epitope** RGPGRAFVTI

Immunogen

Species (MHC) human (A\*0201)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is an A\*0201 epitope.

HXB2 Location gp160 (311-320)

Author Location gp160 (318-327 IIIB)

Epitope RGPGRAFVTI

**Epitope name** LR25

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant (IFA), Montanide (ISA 720), P30, PLG

Species (MHC) mouse (A\*0201)

**Keywords** binding affinity, vaccine-specific epitope characteristics, immunodominance

References Peter et al. 2001

- The stability of peptide binding to HLA-A2.1 was determined for six HLA-A2.1 peptides included in this vaccine study ILKEPVHGV (RT), SLYNTVATL (p17), SLLNATDIAV (gp41) and LLWKGEGAV (RT) all bound with high affinity comparable to a influenza epitope reference (GILGFVFTL), while RGPGRAFVTI and VIYQYMDDL bound with a lower affinity (relative binding activity = 0.01).
- The four high-affinity peptides formed stable complexes with half-lives ranging between 8 and 32 hours, while the low affinity peptides had half lives of less than an hour.
- HLA-A2.1 transgenic mice were immunized with the six HIV-1 peptides and P30, as a universal T-helper epitope, with IFA or Montanide or microspheres as adjuvants.
- All peptides except VIYQYMDDL induced a stong CTL response in Cr-release assays stronger responses were observed when peptides were delivered alone, indicating immunodominance when the combination was used.

HXB2 Location gp160 (311–320)

Author Location gp160 (318-327 IIIB)

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

Species (MHC) human (A2)

References Achour et al. 1996

- Individual was immunized with rec vaccinia gp160 IIIB and boosted with purified gp160.
- Lysis only occurs with IIIB P18 peptide pulsed onto autologous targets; MN, RF, SIMI P18 peptides fail to stimulate CTL.
- Restimulating immune cells from gp160 IIIB vaccinees with MN, RF, or SIMI P18 did not enhance the MN, RF, or SIMI specific CTL response.

HXB2 Location gp160 (311–320)

**Author Location** gp160 (318–327 SIMI)

Epitope MGPKRAFYAT

Immunogen vaccine

Vector/Type: vaccinia prime with gp160 boost Strain: B clade SIMI HIV component: gp160

Species (MHC) human (A2)

References Achour et al. 1996

 Individual was immunized with rec vaccinia gp160 SIMI and boosted with purified recombinant gp160 SIMI.

- P18 MN and RF peptides were able to stimulate the HIVspecific CTL that arose in response to the SIMI vaccination. thus the P18 MN peptide (IGPGRAFYTT) and the P18 RF peptide (KGPGRVIYAT) could cross-react.
- The P18 IIIB peptide does not cross-react (RGPGRAFVTI in the epitope region)
- gp160 SIMI primed immune cells could generate a significantly broader specificity when stimulated with P18 MN or P18RF peptides, but not P18 IIIB.

HXB2 Location gp160 (311-320) Author Location gp120 (311–320) Epitope RGPGRAFVTI Immunogen HIV-1 infection Species (MHC) human (A2)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.

HXB2 Location gp160 (311-320) Author Location gp160 (311–320)

**Epitope** RGPGRAFVTI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, HLA binding

**Keywords** acute/early infection, optimal epitope

References Altfeld et al. 2005

• The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was recognized both in acute and chronic infection, but slightly more frequently in chronic infection.

HXB2 Location gp160 (311-320) Author Location gp160 (318–327 IIIB) Epitope RGPGRAFVTI

Immunogen vaccine

HIV component: V3

Species (MHC) mouse (D)

References Nehete et al. 1995

- · RGPGRAFVTI was defined as the optimal peptide for vaccination, out of RIQRGPGRAFVTIGK.
- This peptide, in a carrier-free form in Freund's adjuvant, could stimulate Env specific CTL in BALB/c mice.

HXB2 Location gp160 (311-320) **Author Location** gp160 (318–327 IIIB) Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

HIV component: V3

**Species (MHC)** mouse (D<sup>d</sup>)

Keywords dendritic cells

References Takahashi et al. 1993

 Successful priming with vaccination of peptide pulsed splenic dendritic cells.

HXB2 Location gp160 (311-320)

Author Location gp160 (318–327 IIIB)

**Epitope** RGPGRAFVTI Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

HIV component: V3

Species (MHC) mouse (D<sup>d</sup>)

References Takahashi et al. 1996

- Exposure of CD8+ CTL to free peptide corresponding to the epitope results in strong inhibition of the CTL response to targets presensitized with the same peptide.
- The authors propose this is due to a "self-veto", where the CTL is inactivated by a CD8+ cell carrying the appropriate peptide-MHC complex.

HXB2 Location gp160 (311–320)

**Author Location** gp160

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: vaccinia

**Species (MHC)** mouse (H-2<sup>d17</sup>)

References Hanke et al. 1998a

- MVA is an attenuated vaccinia that can not replicate in mammalian cells - strings of CTL epitopes were delivered and expressed in a MVA DNA vector.
- INFγ and CTL activity were induced after a single vaccination.
- An MVA boost enhanced the response.

HXB2 Location gp160 (311-320)

**Author Location** gp160

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: DNA, vaccinia HIV compo-

nent: Env Adjuvant: IL-12

Species (MHC) mouse (H-2<sup>d</sup>)

References Gherardi et al. 2000

- Induction of HIV-1 specific CD8 gamma IFN secreting cells was enhanced when IL-12 and Env were given together in a prime, followed by a VV expressing Env boost.
- Vector/Type: peptide Strain: B clade IIIB If IL-12 was also delivered as a boost from the viral vector, impairment of the IL-12 effects was noted, indicating that the vaccination schedule can be a critical parameter for success with DNA and vaccinia vectors used in combination with immunomodulators.
  - The negative effect observed when IL-12 was delivered with the boost involved nitric oxide.

**HXB2 Location** gp160 (311–320)

**Author Location** Env

**Epitope** RGPGRAFVTI

Immunogen vaccine

HIV component: gp160, Rev Adjuvant: IL-12, IL-15, IL-2

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1

References Xin et al. 1999

- A study of the DNA vaccine pCMV160IIIB/REV with IL-15 and IL-2 or IL-12 expression plasmids.
- · Intranasal immunization of BALB/c mice with HIV DNA and IL-15 plasmid induced increased Th1 and CTL responses.
- Co-administration of IL-15 with IL-12 or IL-2 plasmids did not alter the effect of IL-15.
- Both the CTL (peptide pulsed targets) and DTH response (injection of peptide into footpad) to this peptide was monitored.
- The Ab response to NNTRKSIRIQRGPGRAFVTIGKIGN was monitored, and IL-15 co-administration resulted in a decrease in the IgG1/IgG2a ratio.

HXB2 Location gp160 (311-320)

Author Location Env

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: vaccinia, Sindbis HIV component: V3

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Villacres & Bergmann 1999

- HIV-1 epitope p18 was expressed in two different vaccine vectors and the CTL response was compared in BALB/c mice.
- Class I tetramer staining showed that up to 13% of the CD8+ splenocytes were p18 specific in the acute response using vaccinia, only 4% using Sindbis.
- vp18 had more gamma IFN secreting splenocytes and activated CD4+ and CD8+ T cells.
- The overall decline in CD8+ T cells in the transition into memory was 2-3 fold for both vectors.
- · Sindbis virus recombinants induced protective memory cytotoxic T cells, although reduced quantitatively, without vaccinia associated inflamation and replication.

HXB2 Location gp160 (311-320)

**Author Location** Env (318–327)

Epitope RGPGRAFVTI

Immunogen

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords epitope processing, immunodominance

References Lopez et al. 2000

- A series of protease and proteasome inhibitors was used to identify elements of the processing pathway of this epitope, called p18, both from within Env and from within a chimeric hepatitis B protein which allows proper processing.
- · Lactacystin, a proteasome inhibitor, partially inhibits endogenous processing of p18 epitope suggesting both a proteasome pathway and an additional pathway can be used.
- Both TAP dependent and TAP-independent pathways can be
- 1,10-phenanthrolin (metallopeptidases inhibitor) blocks epitope presentation demonstrating metalloproteinase processing in the Tap-dependent pathway.
- The Tap-independent pathway does not involve processing by metalloproteinases.

Vector/Type: DNA Strain: B clade IIIB • This epitope is immunodominant in mice, and is presented by multiple human HLA alleles - it as been suggested that the high processing efficiency of this epitope might result in poor presentation of co-expressed epitopes.

HXB2 Location gp160 (311-320)

Author Location gp120

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: vaccinia

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hanke et al. 1998a; Hanke et al. 1998b

- This murine epitope was incorporated into a vaccine of CTL epitopes expressed together including 20 HIV epitopes recognized by humans from 12 HLA types, one murine HIV epitope and three macaque HIV epitopes, delivered in a vaccinia virus Ankara (VVA) construct.
- The murine vaccination was more effective at generating CTL when given i.v. rather than i.m.

**HXB2 Location** gp160 (311–320)

Author Location gp160 (318–327 IIIB)

Epitope RGPGRAFVTI

Immunogen vaccine

peptide HIV component: Vector/Type: CD4BS, HPG30, V3 Adjuvant: IL-12

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hamajima et al. 1997

- B cell epitope HGP-30 also serves as a CTL epitope.
- Vaccine combined HGP-30, V3 loop peptide variants, and CD4 binding site peptide.
- IL-12 expression plasmid included with the vaccination enhanced the CTL response.

HXB2 Location gp160 (311-320)

Author Location gp160 (318-327 IIIB)

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB

HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1, Th2

References Arai et al. 2000

· Low-dosage 8 Br-cAMP given in combination with a DNA vaccine to BALB/c mice increased IgG and sIgA levels, and enhanced Th1, Th2 and CTL activity - the adjuvant activity may be mediated by activation of the CMV promotor in the DNA vaccine.

HXB2 Location gp160 (311–320)

Author Location gp120 (318–327 IIIB)

**Epitope** RGPGRAFVTI

Immunogen vaccine

Vector/Type: fusion protein with anthrax delivery domain HIV component: gp120

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Goletz et al. 1997

Anthrax lethal toxin can deliver proteins to the cytosol of eukaryotic cells.

 A fusion protein linking the delivery domain of the anthrax protein to gp120 achieved cellular uptake, and gp120 was processed allowing presentation of this V3 epitope to CTL in vitro.

HXB2 Location gp160 (311–320) Author Location gp160 (318–327 IIIB)

**Epitope** RGPGRAFVTI **Epitope name** I-10

Immunogen in vitro stimulation or selection

Species (MHC) mouse (H-2<sup>d</sup>)

References Takahashi et al. 2001

- Pre-incubation of HIV-1 (IIIB) gp160 specific CTL with peptide without APCs reduced cytolytic activity 3.5 fold and induced peptide concentration dependent IL-2 unresponsiveness that might be due to IL-2Rbeta down regulation.
- An enhanced cytolytic activity was observed by addition of anti-IFN-gamma, TNF-alpha or MIP-1beta to I-10 suppressed CTLs.

HXB2 Location gp160 (311–320) Author Location gp160 (IIIB) Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>) **Keywords** Th1, Th2 **References** Shirai *et al.* 2001

Helicobacter pylori induces Th1 responses early, but predominantly Th2 responses later in infection (at 6 weeks) – differentiation of HIV-1 gp160 CD4+ help and CD8+ CTL effector cells in response to HIV gp160-vaccinia vaccination is impaired in BALB/c mice infected with H. pylori.

HXB2 Location gp160 (311–320) Author Location gp120 (V3) (IIIB) Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: influenza Strain: B clade IIIB HIV component: V3

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type Intracellular cytokine staining, Chromium-release assay

**Keywords** genital and mucosal immunity, memory cells, vaccine antigen design

References Garulli et al. 2004

BALB/c mice were transiently infected vaginally with a recombinant influenza virus expressing an HIV CTL V3 epitope.
Infection was promoted by prior progesterone treatment. This
vaccination induced long-term cellular T-cell responses in mice.
Responses were induced at both local mucosal and systemic
sites against both influenza and V3 epitopes. Intranasal vaccination also resulted in T-cell responses in distant mucosal
tissues.

HXB2 Location gp160 (311–320) Author Location gp120 (318–327 IIIB) Epitope RGPGRAFVTI Immunogen vaccine Vector/Type: vaccinia Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>p</sup>, H-2<sup>u</sup>)

References Shirai et al. 1997

- Three class I MHC, H-2<sup>d,p,u</sup>, that differ in sequence and serology, cross-present this peptide to T cells of each of the other haplotypes.
- The amino acids R, F, and I are each critical for strong CTL activity with all three MHC molecules.

HXB2 Location gp160 (311–320) Author Location Env (89.6) Epitope IGPGRARYAR Immunogen vaccine

Vector/Type: vaccinia Strain: B clade 89.6

HIV component: gp160

**Species (MHC)** mouse (H-2D) **References** Belyakov *et al.* 1998b

- Recombinant modified vaccinia virus Ankara (MVA), an attenuated vaccina which has lost the ability to replicate in mammalian cells, was used as the live vector for this vaccine study.
- A single intrarectal mucosal immunization resulted in long lasting mucosal CTL responses and production of proinflammatory cytokines in mucosal sites, indicating that MVA was as effective in inducing mucosal CTL as replicating recombinant vaccinia.

HXB2 Location gp160 (311–320) Author Location Env (IIIB) Epitope IGPGRARYAR

Immunogen vaccine

*Vector/Type:* peptide *Strain:* B clade IIIB *HIV component:* V3

**Species (MHC)** mouse (H-2D) **References** Belyakov *et al.* 1998a

 HIV protection and mucosal CTL response was studied – an HIV peptide immunogen could protect against gp160 expressing vaccinia in a murine intrarectal challenge system in which neutralizing Abs did not play a role, demonstrating mucosal

CTL at the site of exposure can be protective.

HXB2 Location gp160 (311–320) Author Location gp120 (MN) Epitope IGPGRAFYTT Immunogen vaccine

Vector/Type: B. abortus complex

**Species (MHC)** mouse (H-2D<sup>d</sup>) **References** Lapham *et al.* 1996

B. abortus-peptide conjugate induced a virus-specific CTL response in CD4+ lymphocyte depleted mice.

HXB2 Location gp160 (311–320) Author Location gp160 (IIIB) Epitope RGPGRAFVTI Immunogen vaccine

Vector/Type: non-replicating adenovirus Strain: B clade IIIB HIV component: Env,

Rev

**Species (MHC)** mouse (H-2D<sup>d</sup>) **References** Bruce *et al.* 1999

- A good HIV-1 Env immune response using non-replicating adenovirus vectors in BALB/c mice is dependent upon the presence of the stimulatory tat/rev 5'splice-donor site sequence and the presence of Rev.
- Administration of monocistronic RAd501 expressing env and RAd46 expressing rev resulted in a positive CTL response, but required two immunizations for a CTL response comparable to that induced by the bicistronic virus RAd142.
- Administration of RAd501 alone gave a low CTL response, but no humoral response, suggesting a lower level of antigen may be required to stimulate CTL.

HXB2 Location gp160 (311-320) Author Location gp120 (MN) Epitope IGPGRAFYTT Immunogen vaccine

Vector/Type: B. abortus complex

**Species (MHC)** mouse (H-2D<sup>d</sup>)

References Lapham et al. 1996

• B. abortus-peptide conjugate induced a virus-specific CTL response in CD4+ lymphocyte depleted mice.

HXB2 Location gp160 (311-320) Author Location gp160 (318–327 IIIB)

Epitope RGPGRAFVTI

Immunogen peptide-HLA interaction

**Species (MHC)** mouse (H-2D<sup>d</sup>) References Takeshita et al. 1995

• XGPXRXXXXI are critical for binding, consistent with H-2D<sup>d</sup> motif XGPX(RKH)XXX(X)(LIF)

HXB2 Location gp160 (311-320)

**Author Location** Env

Epitope RGPGRAFTVTI

Immunogen vaccine

Vector/Type: DNA HIV component: V3

Species (MHC) mouse (H-2D<sup>d</sup>)

References Hanke & McMichael 1999; Hanke et al. 1999

- · Vaccinated mice elicited a CTL response to a gene gundelivered multiepitope vaccine to two epitopes studied that are known to elicit CTL in mice: SYIPSAEKI from Plasmodium berghei and RGPGRAFTVTI from HIV-1 Env.
- · Different vaccination protocols were tested and it was found that a gene gun mediated delivery followed by an MVA boost was as good as i. m. immunization followed by a MVA boost this is advantageous as gene gun delivery requires far less DNA than i.m. DNA priming.
- CTL activity was high (60% 70% specific lysis at effector target) when vaccinated with a single gene gun immunization and an MVA boost, and improved with two gene gun vaccinations.

**HXB2 Location** gp160 (311–320)

**Author Location** Env (IIIB)

**Epitope** RGPGRAFVTI

Epitope name I-10

Immunogen in vitro stimulation or selection

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Keywords epitope processing, immunodominance

References Nakagawa et al. 2000

- The CTL line LINE-IIIB was generated by repetitive restimulation of BALB/c spleen cells with vSC-25, IIIB gp160expressing vaccinia.
- RGPGRAFVTI represents the active minimal epitope within the previously described immunodominant epitope P18IIIB (RIQRGPGRAFVTIGK, gp160(308-322)
- External processing of P18IIIB results in the removal of the 2 C-terminal residues (GK) of I-10 by ACE (angiotensin-1converting-enzyme) in sera to produce I-10, and this processing is essential for target cell presentation of RIQRGP-GRAFVTIGK.

HXB2 Location gp160 (311–320)

**Author Location** Env (IIIB)

Epitope RGPGRAFVTI

Epitope name p18-I10

Immunogen vaccine

Vector/Type: vaccinia, vesicular stomatitis virus (VSV) Strain: B clade HXB2, B clade IIIB HIV component: Env, Gag

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Keywords immunodominance

References Haglund et al. 2002a

- Different HIV strains were used for different regions: Env IIIB, Gag HXB2
- BALB/c mice were vaccinated with rec vesicular stomatitis virus (rVSV) expressing either HIV-1 Gag, Env, or both, and compared to using rec Env and Gag in vaccinia virus (rVVs). The primary response was determined by cell lysis, cytokine production and tetramer staining.
- Primary CTL responses to the immunodominant Env (RGP-GRAFVTI) epitope peaked 5-7 days after intraperitoneal vaccination with Env-rVSV, 40% of the CD8+ cells were tetramer positive, and this response was 6-fold higher than the response to Env-rVV.
- · Vaccinating with GagEnv-rVSV carrying both Gag and Env allowed recognition of both HIV-1 proteins, but at reduced levels compared to either Gag-rVSV or Env-rVSV alone.
- Intranasal immunization with Env-rVSV yielded CTL responses that were strong but reduced compared to an intraperitoneal route.

HXB2 Location gp160 (311–320)

Author Location Env (IIIB)

Epitope RGPGRAFVTI

Epitope name p18-I10

Subtype B

Immunogen vaccine

Vector/Type: vaccinia, vesicular stomatitis virus (VSV) Strain: B clade HXB2 HIV

component: Env, Gag

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Keywords immunodominance

References Haglund et al. 2002b

- Different HIV strains were used for different regions: Env IIIB, Gag HXB2
- · BALB/c mice were vaccinated with rec vesicular stomatitis virus (rVSV) expressing either HIV-1 Gag or Env, or both, and retention of memory responses and recall responses were studied by tetramer staining and IFN-gamma production.

- Seven months after vaccination with Env-rVSV, 6% of the CD8+ cells were tetramer positive for the immunodominant Env epitope; these cells had a memory phenotype, CD44-Hi positive.
- Env in rec vaccinia virus (Env-rVV) elicited a strong recall response, with up to 45% to the CD8+ T-cell population tetramer positive and activated (expressing CD62L-Lo), and capable of IFN-gamma production.
- A prime with Env-rVSV and heterologous boost of Env-rVV gave remarkably high levels of memory cells, with approximately 1/3 of the CD8+ splenocytes being Env specific memory cells 150 days after the boost.
- A Gag-rVSV or EnvGag-rVSV prime and with a heterologous Gag-rVV or EnvGag-rVV boost combination gave 40% tetramer positive CD8+ cells, but the fraction of IFN-gamma producing cells was only about 25%. Still the heterologous vector prime-boost combination showed a profound benefit.
- A HIV-1 protein rVSV prime, rVV boost was a more potent combination than a vector reversal of a rVV prime and rVSV boost

HXB2 Location gp160 (311–320) Author Location gp120 (V3) (IIIB)

Epitope RGPGRAFVTI

Immunogen vaccine

*Vector/Type:* peptide *Strain:* B clade IIIB *HIV component:* V3 *Adjuvant:* Cholera toxin (CT), GM-CSF, IL-12, IL-18, IL-1 $\alpha$ 

**Species (MHC)** mouse (H-2D<sup>d</sup>) **References** Staats *et al.* 2001

- Cholera toxin (CT) is a potent adjuvant used in animal studies that is not safe in humans, so combinations of cytokins were used in nasal immunization of BALB/c mice V3 peptides to attempt to replace CT as a potent adjuvant.
- Peptide vaccine induced CTL activity was significantly increased by IL-1alpha, IL-18, and GMCSF given alone as adjuvant, but CT gave more potent CTL activity than any single cytokine.
- Combinations of cytokins could be more potent that CT as an adjuvant. The highest tetramer binding of H-2Dd peptidespecific PBMC after nasal immunization was observed with IL-1alpha plus IL-18 as adjuvant.
- Nasal immunization with HIV peptide in the presence of IL-1alpha, IL-12 and GM- CSF induced IFN-gamma-secreting cells in the cervical lymph node, the lung and the spleen, and was associated with upregulation of MHC class II and B7.1 on nonlymphocytes in NALT/nasal mucosal cells.
- Consistent results were obtained for the IIIB and the MN peptides.

**HXB2 Location** gp160 (311–320)

Author Location gp160 (318–327 IIIB)

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: DNA prime with vaccinia boost Strain: B clade IIIB HIV component: gp160 Adjuvant: beta-glucan lentinan, IL-2/Ig, liposome, PLG

**Species (MHC)** mouse (H-2D<sup>d</sup>) **Keywords** immunodominance

References Wierzbicki et al. 2002

• BALB/c mice were give an oral immunization with (PLG)-encapsulated plasmid DNA expressing gp160 and a boost of rec gp160 vaccinia vectors (rVV) with addition of murine IL-2/Ig plasmid or lentinan-associated liposomes. Lentinan increased CTL activity as measured by Cr-release assays against the immunodominant epitope RGPGRAFVTI, but didn't alter Ab responses. IL-2/Ig increased both type I and II activities, and increased Env specific CTL and Abs. Administration of liposomes and PLG microparticles with adjuvants facilitated gastrointestinal uptake.

HXB2 Location gp160 (311–320)

Author Location gp120 (LAI)

Epitope RGPGRAFVTI

**Epitope name** P18

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI HIV component: Gag, gp120 Adjuvant: CpG immunostimulatory sequence (ISS)

**Species (MHC)** mouse (H-2D<sup>d</sup>) **References** Horner *et al.* 2001

 Immunostimulatory sequences (ISS), also known as CpG motifs, stimulate innate immunity and enhance vaccine-specific immune responses.

- Intranasal immunization (i.n.) of BALB/c mice was more effective than intradermal (i.d.), and immunization with a gp120-ISS conjugate was more potent than immunizing with gp120 and separate ISS molecule increased IgG1, IgG2a, IFN-gamma, MIP1-alpha and MIP1-beta production was observed, and only i.n. immunization gave IgA responses.
- The highest mucosal CTL activity in both the Lamina Propria and the Peyer's Patch was observed following intranasal delivery with the gp120/ISS conjugate.
- Cytokine, chemokine and CTL responses following gp120/ISS conjugate vaccination were CD4+ T-cell independent; gp120 specific antibodies were dependent on helper T cells.

**HXB2 Location** gp160 (311–320)

Author Location gp160 (V3) (IIIB)

Epitope RGPGRAFVTI

Epitope name I10

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Keywords acute/early infection

**References** Takahashi *et al.* 2002

- During acute infection, high doses of virus result in "clonal exhaustion", a depletion of antigen specific T-cells.
- Recently stimulated CTL from BALB/c mice vaccinated with gp160-vaccinia showed a dose- and time-dependent induction of apoptosis when stimulated with antigenic peptide or H-2Dd/peptide tetramers.
- Restimulated CTL showed an upregulation of CD3-chain phosphorylation in comparison to cells stimulated with target cells, indicative of TCR-mediated apoptosis. Furthermore, apoptosis was inhibited by cyclosporin A and U0126, a mitogen activated

kinase inhibitor specific for the ERK1/ERK2 MAPK kinase pathway, and a caspase 3 inhibitor.

HXB2 Location gp160 (311–320)
Author Location gp160 (V3) (MN)
Epitope IGPGRAFYAT
Epitope name MNT10
Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

Species (MHC) mouse (H-2D<sup>d</sup>)

Keywords acute/early infection

References Takahashi *et al.* 2002

- During acute infection, high doses of virus result in "clonal exhaustion", a depletion of antigen specific T-cells.
- Recently stimulated CTL from BALB/c mice vaccinated with gp160-vaccinia showed a dose- and time-dependent induction of apoptosis when stimulated with antigenic peptide or H-2Dd/peptide tetramers.
- Restimulated CTL showed an upregulation of CD3-chain phosphorylation in comparison to cells stimulated with target cells, indicative of TCR-mediated apoptosis. Furthermore, apoptosis was inhibited by cyclosporin A and U0126, a mitogen activated kinase inhibitor specific for the ERK1/ERK2 MAPK kinase pathway, and a caspase 3 inhibitor.

HXB2 Location gp160 (311–320) Author Location gp160 (V3) (HIV-IIIB)

Epitope RGPGRAFVTI Epitope name P18-I10 Subtype B

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160 Adjuvant: IL-15,

IL-2

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Donor MHC H-2d

Assay type Cytokine production, Tetramer binding, Chromium-release assay

References Oh et al. 2003a

• IL-2 and IL-15 in vaccinia constructs were given with an HIV gp160 vaccinia vaccine to BALB/c mice. Both IL-2 and IL-15 induced strong and long-lasting antibody responses. Short-term CTL responses against HIV gp120 were enhanced by IL-2, but IL-15 enhanced both immediate CD8+ T cell responses and CD8+ T memory cells.

HXB2 Location gp160 (311–320) Author Location gp160 (IIIB)

Epitope RGPGRAFVTI

Epitope name P18-I10 Subtype B

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: V3 Adjuvant: B7, ICAM,

LFA-3

Species (MHC) mouse (H-2D<sup>d</sup>)

Donor MHC H-2d

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Tetramer binding

## References Oh et al. 2003b

BALB/c mice were vaccinated with T-cell depleted splenocytes
pulsed with peptides given in combination with immunostimulatory molecules B7, ICAM or LFA expressed in a recombinant
pox virus. Increasing antigen gave an increased frequency of
CD8+ T-cells, but the co-stimulatory molecules increased the
avidity of the response.

HXB2 Location gp160 (311-320)

**Author Location** (89.6)

Epitope IGPGRAFYAR

**Subtype** B **Immunogen** vaccine

Vector/Type: DNA HIV component: gp120 Adjuvant: Flex, a dendric cell growth factor

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Donor MHC H-2d

Assay type Intracellular cytokine staining

**Keywords** dendritic cells **References** Sailaja *et al.* 2003

- BALB/c mice were given a DNA vaccine that contained gp120 DNA covalently attached to the extracellular domain of the Fms-like tyrosine kinase receptor-3 ligand (FLex), a dendritic cell growth factor.
- Mice vaccinated i.m. with the FLex:gp120 chimeric gene gave a DC expansion similar to native Flex protein.
- gp120-specific stable CD8+ T-cell responses lasted 114 days after a prime/boost, and were observed in the presence and absence of Flex-DNA-induced dendritic cell (DC) expansion; strong Ab responses required DC expansion.

**HXB2 Location** gp160 (311–320)

Author Location gp120 (V3)

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: herpes simplex virus type-1 (HSV-1) amplicon HIV component: gp120

Species (MHC) mouse (H-2D<sup>d</sup>)

Donor MHC H-2d

Assay type Tetramer binding, JAM cytotoxicity assay

Keywords kinetics, memory cells

References Wang 2003

- Prime-boost combinations of gp120 combined with herpes simplex virus type-1 (HSV-1) amplicon particles, or gp120 in naked amplicon plasmid DNA, were compared in BLAB/c mice. Plasmid prime with particle boosts gave the strong primary (2 weeks) and memory responses (4 months).
- CD8+ T-cells reached their peak 8-28 days after the initial amplicon delivery.

HXB2 Location gp160 (311-320)

Author Location gp120 (V3)

Epitope RGPGRAFVTI

Epitope name P18-I10

Immunogen vaccine

Vector/Type: peptide, vaccinia Strain: B clade 89.6, B clade IIIB HIV component: gp160ΔV3 Adjuvant: Cholera toxin (CT), E. coli mutant heat labile enterotoxin (LT-R72),

Incomplete Freund's Adjuvant (IFA), CpG immunostimulatory sequence (ISS)

Species (MHC) mouse (H-2D<sup>d</sup>)

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay

Keywords dendritic cells, Th1, Th2, genital and mucosal

immunity

References Belyakov et al. 2004

• Transcutaneous immunisation (TCI) of BALB/c mice induced adjuvant-dependent HIV-1 specific CTL responses in the spleen and the gut mucosa that resulted in protection against mucosal challenge against a recombinant vaccina virus carrying HIV-1 env. Activated DCs from skin were shown to migrate to immune-inductive sites in gut mucosa and to present antigen directly to resident lymphocytes.

HXB2 Location gp160 (311-320)

Author Location gp120 (V3)

**Epitope** RGPGRAFVTI

Subtype B

Immunogen vaccine

Vector/Type: herpes simplex virus type-1 (HSV-1) amplicon Strain: B clade LAI, B clade MN HIV component: gp120

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Assay type CD8 T-cell Elispot - IFNy, Tetramer binding,

JAM cytotoxicity assay Keywords vaccine antigen design

References Hocknell et al. 2002

· BALB/c mice were immunized with HSV amplicons containing HIV-1 gp120. Helper virus free HSV-1 amplicon particles are capable of inducing potent cytoxic CD8+ T-cell and humoral immune responses to the HIV-1 antigen in mice. Previous infection with wild-type HSV-1 reduces amplicon-induced cellular immune responses to HIV gp120 modestly (40-60%), but severally reduced B-cell responses. The route of vaccination impacted the nature and level of the responses (i.m., i.d., and i. p.).

HXB2 Location gp160 (311-320)

Author Location gp120

Epitope RGPGRAFVTI

Subtype B

Immunogen vaccine

Vector/Type: DNA, polyepitope Strain: B clade MN HIV component: gp120, Protease, RT Adjuvant: Incomplete Freund's Adjuvant (IFA)

**Species (MHC)** humanized mouse (H-2D<sup>d</sup>)

Assay type CD8 T-cell Elispot - IFNγ

Keywords epitope processing, vaccine-specific epitope characteristics, immunodominance, immunotherapy

References Isaguliants et al. 2004

• Immunization of HLA-A\*0201-transgenic mice with synthetic genes encoding clusters of human A\*0201 CTL epitopes located at the sites of drug resistance mutations, induced RTspecific cellular responses indicating the immunogenicity of these constructs. This vaccine strategy may be a first step towards a therapeutic vaccine against drug-resistant strains. This epitope was included as a mouse marker for a CD8+ T-cell response.

HXB2 Location gp160 (311–320)

**Author Location** 

Epitope RGPGRAFVTI

Epitope name R10I

Immunogen vaccine

Vector/Type: DNA, virus-like particle (VLP), polyepitope HIV component: Gag, p24 Gag,

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Assay type Cytokine production, Chromium-release as-

Keywords epitope processing, vaccine-specific epitope characteristics, immunodominance

References Wild et al. 2004

- · A codon optimized gag DNA vaccine was compared to a myristylation defective gag and p24 alone, both of which lack signals for secretion from transfected cells. Gag-derived immunogens that were secreted as VLPs and those that remained intracellular (p24) each produced strong CTL responses, and neither the size of antigen nor cellular trafficking and localization significantly influenced the strength of humoral and cellular immune activation. The formation and release of VLPs was not essential for eliciting strong CTL. BALB/c mice were given the DNA vaccine by i.m. administration of plasmid DNA for the prime and boost.
- Linking the region encoding the V3 immunodominant epitope to the gag gene did not diminish the response to the Gag p24 epitope A9I, but did enable a response to the V3 epitope.
- Minigenes were made incorporating just 1 epitope, minitopes, carrying 1 of 3 murine class I epitopes linked to the Ad2-E3 protein-derived signal peptide to allow access of the epitope to the ER. Weak induction of cellular immune responses was observed, in contrast to the complex polyprotein.

HXB2 Location gp160 (311–320)

**Author Location** Env (318–327)

Epitope RGPGRAFVTI

Epitope name R10I

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Donor MHC H-2D

Assay type Intracellular cytokine staining

Keywords vaccine antigen design

References Samino et al. 2004

• The endogenous processing of the HIV-1 envelope glycoprotein generates several different natural peptidic species presented by the H-2D molecule in infected cells, the 9-, 10-, and 11-mer peptides, GPGRAFVTI, RGPGRAFVTI, and QRGP-GRAFVTI. CTL with the same antigenicity could recognize all three forms. The complexity of the binding peptides suggests naturally processed proteins could provide more variety as antigens, stimualting more robust and diverse CTL responses.

HXB2 Location gp160 (311–320)

Author Location p18 (IIIB)

Epitope RGPGRAFVTI

Subtype B

Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade IIIB HIV component: gp120

Adjuvant: IL-12

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Donor MHC H-2D

Assay type Cytokine production, proliferation, CD8 Tcell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ,

Chromium-release assay

Keywords vaccine-induced epitopes, memory cells, char-

acterizing CD8+ T cells

References Seaman et al. 2004

 Delivery of plasmid IL-12 on day 10 after immunization of mice with an HIV-1 gp120 DNA vaccine resulted in expansion of gp120-specific CD8+ T-cells but had no effect on antigenspecific CD4+ T-cells and antibody responses. gp120-specific CD8+ T-cells were shown to primarly be effector memory and not central memory T-cells and did not expand following gp120 boost immunization.

HXB2 Location gp160 (311–320)

Author Location gp160 (318–327 IIIB)

**Epitope** RGPGRAFVTI **Immunogen** vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: Env, Nef

**Species (MHC)** mouse (L<sup>d</sup>)

References Tobery & Siliciano 1997

- An HIV-1 Env vaccine was targeted for rapid cytoplasmic degradation.
- The rapidly degraded form rapidly stimulated CTL to this peptide, faster than the normal vaccinia-env.
- The rapidly degraded form also stimulated greater specific CTL lysis and higher CTLp frequencies than normal Env.
- Similar results were obtained for a Nef protein designed for rapid degradation.

**HXB2 Location** gp160 (311–320)

 $\textbf{Author Location} \ \ gp160 \ (318\text{--}327 \ IIIB)$ 

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: DNA prime with peptide boost Strain: B clade IIIB HIV component:

CD4BS, gp160, HPG30, V3

Species (MHC) macaque

References Okuda et al. 1997

• Murine BALB/c (H-2<sup>d</sup>) and macaque both showed highest level of CTL vaccine response when a DNA vaccine was boosted with a peptide including four peptide subtypes of the V3 region, HPG-30 and a fragment of the CD4 binding region.

HXB2 Location gp160 (311–320)

Author Location gp120 (318–327)

Epitope RGPGRAFVTI

Immunogen HIV-1 infection

Species (MHC) human

References Kmieciak et al. 1998b

- Increased CTL response to cells expressing a VV construct ΔV3 mutant compared with a full-length env gene product.
- This epitope doesn't have A2 anchors, but has features that confer promiscuous A2 binding, which may relate to the inhibitory effect seen in this paper.

HXB2 Location gp160 (311-320)

**Author Location** Env (IIIB)

Epitope RGPGRAFVTI

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB HIV component: gp160, Rev Adjuvant:

MIP-1 $\alpha$ 

Species (MHC) mouse

References Lu et al. 1999

- MIP-1alpha co-inoculation increased IgG1/IgG2a ratio Thelper type 1 response.
- A MIP-1 alpha expression plasmid increased the CTL response to this DNA vaccine, as well as the T help response, presumably by the MIP-1 alpha interacting with T lymphocytes and macrophages.

HXB2 Location gp160 (311–320)

**Author Location** 

Epitope RGPGRAFVTI

Epitope name P18

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade BH10 HIV component: gp120 Adjuvant: GM-CSF

Species (MHC) mouse

References Barouch et al. 2002

- gp120 encoding DNA co-injected with a plasmid carrying GM-CSF gave meager CD4+ T-cell responses in BALB/c mice relative to the enhanced response to bicistronic gp120 and GM-CSF cloned into the same vector and expressed from the same promoter.
- Both mono and bicistronic DNA vaccines induced similar CTL responses directed against the H-2Dd restricted P18 peptide RGPGRAFTVTI in murine splenocytes despite the greatly enhanced proliferative responses.

**HXB2 Location** gp160 (311–320)

Author Location gp120 (313-322 BRU)

Epitope RGPGRAFVTI

Epitope name Pep 09

**Subtype** B, C **Immunogen** vaccine

Vector/Type: DNA Strain: B clade BRU

HIV component: gp160, Rev, Tat

Species (MHC) mouse

Keywords subtype comparisons, Th1

References Arora & Seth 2001

- Plasmid DNA encoding gp160, tat, rev was given i.m. to immunize BALB/c mice.
- Vaccine-induced CTL activity produced a low degree of cell lysis of V3-peptide pulsed target cells, using a B (RGP-GRAFVTI) or C (RIGGPGQTFYATG) clade V3 peptides. Th1 proliferative T-cell responses were observed, and weak Ab responses.

HXB2 Location gp160 (311-320) Author Location Env (IIIB) Epitope RGPGRAFVTI Epitope name 10 Env Subtype B Immunogen vaccine

> Vector/Type: influenza prime with vaccinia boost Strain: B clade IIIB HIV compo-

nent: gp160

Species (MHC) mouse Donor MHC H-2d

> Assay type Cytokine production, proliferation, CD8 Tcell Elispot - IFNγ

**Keywords** Th1, Th2, genital and mucosal immunity References Gherardi et al. 2003

- Mice were intranasally primed with a recombinant influenza virus A vector that carries HIV-1 Env inserted into its hemagglutinin protein. Boosting was performed intranasally with either influenza-Env or intraperitoneally with two vaccinia virus recombinants expressing the Env protein, VVenv and MVAenv.
- · Peritoneal heterologous immunization with VVenv induced a 60-fold higher CD8+ IFN-gamma T cell responses than homologous influenza prime-boost. The intraperitoneal MVAenv boost response was greater than the VVenv boost in the spleen and genital lymph nodes, while the VVenv response gave the highest boost with the intranasal route.
- Mice with increased CD8+-T-cell responses also had a higher Th1/Th2 ratio, indicated by the cytokine secretion profile and the IgG2a/IgG1 ratio.

HXB2 Location gp160 (311-320)

Author Location gp160

Epitope RGPGRAFVTI

Epitope name P18-I10

Subtype B

Immunogen vaccine

Vector/Type: vaccinia with H1 influenza HA gene cassette Strain: B clade IIIB HIV

component: gp160

Species (MHC) mouse

**Assay type** Chromium-release assay

Keywords genital and mucosal immunity

References Kuribayashi et al. 2004

• The intraepithelial compartment of the intestinal mucosa is shown to be a major site for preventing virus spread by thymus-derived CD8alphaß-positive Ag specific CTLs and CD8alpha, alpha+gamma, delta cells, which regulate virus spread in a P18-I10 vaccinia vector mouse infection model.

HXB2 Location gp160 (312–320)

Author Location gp120 (V3) (IIIB)

Epitope GPGRAFVTI

Subtype B

Immunogen vaccine

Vector/Type: fowlpoxvirus Strain: B clade BRVA, B clade IIIB, B clade JY1, B clade LR150, B clade MN, B clade RF HIV com-

ponent: V3

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords vaccine-specific epitope characteristics, immunodominance

References Vázquez Blomquist et al. 2002

- BALB/c mice were vaccinated with a polyepitope V3 vaccine in a fowlpoxvirus carrying concatonated 15 mer sections of the V3 loops of HIV-1 isolates LR150, JY1, RF, MN, BRVA and IIIB with 5-aa linkers between, fused to the N-term of p64K protein from Neisseria meningitidis.
- Intraperitoneal immunization elicited the strongest V3-specific IFN-gamma response in splenocytes, compared to intravenous and subcutaneous immunization. Intraperitoneal immunization conferred protection in a recombinant vaccinia virus challenge
- The immunodominant response was directed against the IIIB peptide (the IIIB immunizing peptide was SIRIQRGP-GRAFVTI, the peptide used to probe the response by Elispot was GPGRAFVTI).
- Low CTL responses were also detected to the LR150 (SR-GIRIGPGRAILAT) and RF (RKRITMGPGRVYYTT) peptides, no responses were detected to the JY1 (RQSTPIGLGQ-ALYTT), BRVA (RKSITKGPGRVIYAT), or MN (RKRIHIGP-GRAFYTT) peptides.

HXB2 Location gp160 (312–320)

Author Location gp120 (V3)

Epitope GPGRAFVTI

Subtype B Immunogen vaccine

Vector/Type: DNA prime with vaccinia boost, polyepitope, DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade BRVA, B clade IIIB, B clade JY1, B clade LR150, B clade MN, B clade RF HIV com-

ponent: V3 Adjuvant: IFN \u03b7 **Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type Cytokine production, CD8 T-cell Elispot -IFNγ

Keywords Th1

References Gómez et al. 2004

- Priming of mice with DNA-TAB vector, a polyepitope string carrying 8 different V3 loop sequences, followed by a booster with VV-TAB or MVA-TAB, induced humoral responses, as well as a CD8+ T-cell response against V3 epitopes from three different subtype B HIV isolates. The highest values of specific CD8+ T-cell response were achieved when priming with DNA-TAB and a DNA vector expressing IFN-gamma, followed by a MVA-TAB boost. The T-cell response was Th1.
- The eight V3 loops were linked with an A-G-G-G-A sequence. The three peptides that elicited a response were LR150, SRGIRIGPGRAIL; MN, RKRIHIGPGRAFY; and IIIB, SIRIQRGPGRAFVTI. These peptides were located at the beginning, middle and end of the polyepitope, indicating all parts were able to be processed. It is not known if there is an H-2d epitope in the other five V3 loop variants that did not elicit a response.

**HXB2 Location** gp160 (314–322)

**Author Location** gp120 (312–320)

Epitope GRAFVTIGK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*2705)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- One of seven patients responded to this peptide with GzB producing cells and with IFN-gamma producing cells.

**HXB2 Location** gp160 (314–322)

Author Location gp120 (314–322)

Epitope GRAFVTIGK

Immunogen peptide-HLA interaction

Species (MHC) human (B27)

References Jardetzky et al. 1991

• Study of peptide binding to HLA-B27.

HXB2 Location gp160 (321-330)

Author Location gp120 (322–330 HIV-MN)

Epitope EIIGDIRQAY

Epitope name EY10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2501)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- This is a newly defined epitope. Positions 1, 5 and 10 (last)in the epitope had potentially experienced positive selection. qI-IGDIRQAY, dIIGDIRQAh, EIIGnIRQAh and EIIGDIRQAh escape variants were found.

**HXB2 Location** gp160 (337–361)

Author Location gp120 (337–368 LAI)

Epitope KWNNTLKQIDSKLREQFGNNKTIIF

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (CD4+ CTL)

References Johnson et al. 1994a

 CD4+ CTL clones were obtained from an HIV-1 vaccinia-env vaccinee.

**HXB2 Location** gp160 (339–354)

Author Location gp120 (339-361 LAI)

Epitope NNTLKQIDSKLREQFG

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (CD4+ CTL)

References Johnson et al. 1994b

• CD4+ CTL isolated from LAI IIIB gp160 vaccinees.

**HXB2 Location** gp160 (340–348)

Author Location gp120 (346–354 CM243 subtype CRF01)

Epitope RVLKQVTEK
Epitope name E340-348
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A11)

Keywords HIV exposed persistently seronegative

(HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was weakly reactive in HIV+ control study subject 053 who carried HLA-A11.

HXB2 Location gp160 (340–348)

Author Location gp120 (346–354 CM243 subtype CRF01)

Epitope RVLKQVTEK Subtype CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A11)

**Keywords** subtype comparisons **References** Bond *et al.* 2001

- HLA-A11 CRF01 (called subtype E in Bond *et al.*) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This is one of the new A11 epitopes identified through the streamlined EpiMatrix method, and 2/8 tested FSWs recognized it.
- This epitope was not conserved in other subtypes, and exact matches were rare.

**HXB2 Location** gp160 (340–349)

Author Location gp120 (W6.ID)

Epitope NTLKQIVIKL

Immunogen vaccine

Vector/Type: protein Strain: B clade W61D

HIV component: gp120

Species (MHC) chimpanzee (Patr-B\*14)

Keywords immunodominance

References Balla-Jhagjhoorsingh et al. 1999a

 An HIV-1 rgp120 vaccine induced strong humoral and cellular immune response in sibling chimpanzees, but only one of the two made a detectable CTL response to this Patr-B\*14 restricted immunodominant epitope.

HXB2 Location gp160 (341–349)

Author Location gp120 (341–349 HIV-MN)

Epitope TLSQIVTKL

Epitope name TL9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*0201)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- This is a newly defined epitope. Position 4 in the epitope had potentially experienced positive selection. TLSkIVTKL escape variant was found.

**HXB2 Location** gp160 (344–361)

Author Location gp160 (348–366 WEAU)

Epitope QIVEKLREIKQFKNKTIVF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*2902, B\*4403, B\*0801

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

Keywords dynamics, immunodominance, escape, kinet-

ics, characterizing CD8+ T cells, reversion,

viral fitness

References Jones et al. 2004

Primary CD8+ T-cell response to Env, Tat and Gag and the
extent, kinetics and mechanisms of viral escape were examined
in three patients. Rapid escape, within weeks from infection,
from HIV specific CTL responses was observed in all three
patients, but the kinetics and extent of the escape differed
depending on the breadth and co-dominant distribution of CTLmediated pressure. The two patients that rapidly declined had
more focused immunodominant responses, while the single

patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.

- The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance. WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.
- WEAU had a reaction to an eptiope within this peptide, and there was very rapid accumulation of substitutions; variation continued through the last sample collected.

HXB2 Location gp160 (363–376)

Author Location (C consensus)

**Epitope** PSSGGDLEITTHSF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*18)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** gp160 (369–375)

**Author Location** gp120 (374–380 BRU)

Epitope PEIVTHS

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Dadaglio et al. 1991

• Defined through blocking CTL activity, and Env deletions.

HXB2 Location gp160 (374–382)

**Author Location** Env

Epitope HSFNCGGEF

Epitope name 1325

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A02, A03, B08, B51, Cw01, Cw07

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for HSFNCGGEF: 76%

**HXB2 Location** gp160 (375–383) **Author Location** gp120 (379–387 LAI)

**Epitope** SFNCGGEFF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1516)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1516 epitope.

HXB2 Location gp160 (375–383)

Author Location Env

**Epitope SFNCGGEFF** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1516)

**Donor MHC** A\*0202, A\*0301, B\*0702, B\*1516

Country United States.

Keywords escape, acute/early infection

References Bernardin et al. 2005

- Full HIV genomes from nine individuals were analyzed for mutations prior to seroconversion and 7 to 28 days later. It was found that the influence of the host's HLA type was reflected within weeks of infection; a statistically significant number of early nonsynonymous mutations were observed within previously reported CTL epitopes.
- E to G mutation was observed in position 7.

HXB2 Location gp160 (375–383)

**Author Location** gp120 (375–383 IIIB)

Epitope SFTCGGEFF Immunogen HIV-1 infection Species (MHC) human (B15)

Keywords responses in children, mother-to-infant trans-

mission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- An additional variant that gave a positive, though reduced, CTL response: SSTCGGEFF and SFTCGGGFF.
- SFTCGGGVF was an escape mutant.

HXB2 Location gp160 (375–383)

Author Location gp120 (375–383 SF2)

Epitope SFNCGGEFF Immunogen HIV-1 infection Species (MHC) human (B15)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic

infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.

- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B15+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 1/1 group 2, and 0/1 group 3.

**HXB2 Location** gp160 (375–383)

**Author Location** gp120 (375–383)

**Epitope** SFNCGGEFF

Immunogen HIV-1 infection

Species (MHC) human (B15)

**Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$  **Keywords** rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 5 epitopes from this individual that varied over time where the internal mutations became fixed; a dramatic decrease in CTL activity against the wild type epitope was observed as the mutation arose. The sfncRgeff variant residue found at 20 and 47 months postseroconversion.

HXB2 Location gp160 (375–383)

Author Location gp120

**Epitope** SFNCGGEFF

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human (B63)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This is a putative HLA-B63/57/58 epitope containing the B58 supertype binding motif. Borderline significantly more often recognized in subjects who carry B63, but not B57/B58.

**HXB2 Location** gp160 (375–383)

**Author Location** gp120 (375–383 IIIB)

Epitope SFNCGGEFF

Immunogen HIV-1 infection

Species (MHC) human (B63, B15)

References Wilson et al. 1997a

- This is the optimal peptide for two CTL clones that recognize this epitope in the context of two different HLA molecules, Cw4 and B15.
- Predominant form in proviral DNA of the individual with B15 restricted CTL was SFTCGGEFF and this was recognized.
- Recognition of a minor autologous variant (SFNCRGEFF) from the B15 donor was greatly reduced.

HXB2 Location gp160 (375–383)

**Author Location** gp120 (376–383 PV22)

**Epitope** SFNCGGEFF

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0401)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a C\*0401 epitope.

**HXB2 Location** gp160 (375–383)

Author Location gp120

**Epitope** SFNCGGEFF

Subtype A

Immunogen HIV-1 infection

**Species (MHC)** human (Cw\*0401, Cw\*0407)

**Keywords** HIV exposed persistently seronegative (HEPS), cross-presentation by different HLA

References Bird et al. 2002

- 4/123 (2 HIV-1 positive, 2 HEPS) Kenyan female sex workers carried the novel allele HLA Cw\*0407.
- HLA Cw\*0407 did not differ from Cw\*0401 in the region associated with the binding pocket, and Cw\*0407 was shown to cross-present a previously defined Cw\*0401 epitope, SFNCGGEFF (gp120).

**HXB2 Location** gp160 (375–383)

Author Location gp120 (376-383 PV22)

**Epitope** SFNCGGEFF

Immunogen HIV-1 infection

Species (MHC) human (Cw4)

References Johnson et al. 1993

· Conserved epitope.

HXB2 Location gp160 (375–383)

**Author Location** gp120 (376–383 PV22)

Epitope SFNCGGEFF

Immunogen HIV-1 infection

Species (MHC) human (Cw4)

References Wolinsky et al. 1996

• Longitudinal study of epitope variation in vivo.

HXB2 Location gp160 (375–383)

**Author Location** gp120 (376–383)

**Epitope** SFNCGGEFF

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (Cw4)

Keywords HIV exposed persistently seronegative

(HEPS), immunodominance

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused
  on different epitopes with HLA presenting molecules that have
  previously been associated with reduced risk of infection, and
  there was a shift in the response in the HEPS women upon late
  seroconversion to epitopes recognized by the HIV-1 infected
  women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-Cw4 women, 1/2 HEPS and 10/11 HIV-1 infected women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in 6 of the 10/11 responsive HIV-1 infected women, and not in the HEPS case.

HXB2 Location gp160 (375–384)

**Author Location** (C consensus)

**Epitope** SFNCRGEFFY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*29)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- SFNCRGEFFY is an optimal epitope.

**HXB2 Location** gp160 (375–384)

Author Location (B consensus)

**Epitope** SFNCGGEFFY

**Epitope name** SY10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A29)

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

**Assay type** Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
   activation in dying target cells, it was shown that the subset of
   HIV-1-specific CD8+ T cells secreting both IFN-gamma and
   TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1 specific CD8+ T-cell maturation phenotypes and intracellular
   perforin expression was found.
- 1/9 individuals recognized this epitope.

HXB2 Location gp160 (376-383)

Author Location gp120

**Epitope** FNCGGEFF

Immunogen

Species (MHC) human (Cw4)

**References** Rowland-Jones *et al.* 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied - these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive,
- HIV-2 sequence: TNCRGEFL no cross-reactivity Johnson et al. [1993]

HXB2 Location gp160 (376-384)

Author Location gp120 (376–384 IIIB)

**Epitope** FNCGGEFFY Immunogen HIV-1 infection Species (MHC) human (A29)

References Wilson et al. 1997a

- This is the optimal peptide for two CTL clones derived from two different donors.
- · FNCRGEFFY and FNCRGGFFY are major and minor autologous variants in one of the donors, and showed reduced or no stimulatory activity for CTL from the host.
- The IIIB form and the form FNCAGEFFY were present in the other donor, and the CTL line had reduced activity with the FNCAGEFFY form relative to the index peptide.

HXB2 Location gp160 (376-384)

Author Location gp120 (376–384 IIIB)

Epitope PNCGGEFFY Immunogen HIV-1 infection Species (MHC) human (A29)

**Keywords** responses in children, mother-to-infant transmission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- · Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- PNCRGEFFY was an escape variant.

HXB2 Location gp160 (376-384)

Author Location gp120 (376-384 LAI)

**Epitope** FNCGGEFFY

Epitope name E2

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A29)

Keywords HAART, ART

References Mollet et al. 2000

· A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.

- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened – eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+ PBL - but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

HXB2 Location gp160 (376–384)

Author Location gp120

**Epitope** FNCGGEFFY Immunogen HIV-1 infection Species (MHC) human (A29)

Assay type Intracellular cytokine staining

Keywords immunodominance, genital and mucosal immunity

References Kaul et al. 2003

- Predefined immunodominant peptide responses were used to compare CD8+ T cells responses in the blood and cervix of 16 HIV+ Kenyan sex workers. Cervical responses were detected in 8/10 women from whom adequate samples could be obtained. The frequency of the CD8+ T cell response in the genital tract was comparable to the blood, with a trend toward being slightly
- The immunodominant response was to this epitope in the PBMC of 1/16 patients (Kaul et al. 2001, AIDS, 107:1303).

HXB2 Location gp160 (376–384)

Author Location gp120 (376–384)

**Epitope** FNCGGEFFY

**Epitope name** FNC

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Keywords** HAART, ART, supervised treatment interruptions (STI), immunodominance, acute/early

infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- One of the 7/8 study subjects that were HLA B8 recognized this CTL epitope.
- Patient SC19(HLA A11/12, B8/44, Cw06/0701, DR3/7, DR52/53, DQ 2/8) had a CTL response to epitopes FLKEKGGL, GEIYKRWII, ACQGVGGPGHK, AVDLSH-FLK, and FNCGGEFFY that declined during therapy initiated at day 197.

HXB2 Location gp160 (376-384)

Author Location gp160

**Epitope** FNCGGEFFY

Epitope name FNC

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius et al. [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there was no correlation between CTL responses with viral rebound rates, plateau viral loads, or clearance rates.

**HXB2 Location** gp160 (376–387)

Author Location gp120 (381-392 BRU)

Epitope KNCGGEFFYCNS

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Dadaglio et al. 1991

• Defined through blocking CTL activity, and Env deletions.

**HXB2 Location** gp160 (376–387)

**Author Location** Env (379–)

**Epitope** KNCGGEFFYCNS

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A\*0202, A\*0301, B\*0702, B\*1516

**Country** United States.

Keywords escape, acute/early infection

References Bernardin et al. 2005

- Full HIV genomes from nine individuals were analyzed for mutations prior to seroconversion and 7 to 28 days later. It was found that the influence of the host's HLA type was reflected within weeks of infection; a statistically significant number of early nonsynonymous mutations were observed within previously reported CTL epitopes.
- E to G mutation was observed in position 6.

**HXB2 Location** gp160 (377–386)

Author Location gp160 (374–383 SUMA)

Epitope NCGGEFFYCN

Epitope name gp160 NN10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

**Country** United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, acute/early infection, characterizing CD8+ T cells

References Jones et al. 2004

Primary CD8+ T-cell response to Env, Tat and Gag and the
extent, kinetics and mechanisms of viral escape were examined
in three patients. Rapid escape, within weeks from infection,
from HIV specific CTL responses was observed in all three
patients, but the kinetics and extent of the escape differed
depending on the breadth and co-dominant distribution of CTLmediated pressure. The two patients that rapidly declined had

more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.

- The patient SUMA maintained low viral loads and stable CD4
   T cell counts through seven years of follow up. In contrast
   to more rapid progressors, WEAU and BORI, SUMA a broad
   response to 24 epitopes, with little immunodominance. Two
   peptides were somewhat more intensely recognized in acute
   infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

**HXB2 Location** gp160 (377–387)

**Author Location** gp120 (377–387)

**Epitope** NSGGEFFYSNS

Immunogen

Species (MHC) human (A2)

References Hickling et al. 1990

Peptides recognized by class I restricted CTL can bind to class
II

**HXB2 Location** gp160 (383–391)

Author Location gp120 (385–393)

**Epitope** FYCNTTQLF

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

References Ikeda-Moore et al. 1997

• Defined using reverse immunogenetics – 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) – 53 of the 59 peptides bound A\*2402.

• This peptide induced CTL in 1/4 HIV-1 + people tested.

 FYCNTTQLF bound to A\*2402 strongly, the epitope can be processed in a vaccinia construct and presented – two specific CTL clones were obtained.

**HXB2 Location** gp160 (383–391)

Author Location gp160 (380–389 SUMA)

Epitope FYCNTTQLF

**Epitope name** GP160 FF9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, acute/early infection, characterizing CD8+ T cells

References Jones et al. 2004

Primary CD8+ T-cell response to Env, Tat and Gag and the
extent, kinetics and mechanisms of viral escape were examined
in three patients. Rapid escape, within weeks from infection,
from HIV specific CTL responses was observed in all three
patients, but the kinetics and extent of the escape differed
depending on the breadth and co-dominant distribution of CTLmediated pressure. The two patients that rapidly declined had

more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.

- The patient SUMA maintained low viral loads and stable CD4 T cell counts through seven years of follow up. In contrast to more rapid progressors, WEAU and BORI, SUMA a broad response to 24 epitopes, with little immunodominance. Two peptides were somewhat more intensely recognized in acute infection, but this response leveled out early on.
- Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

HXB2 Location gp160 (410-429) **Author Location** gp120 (410–429 PV22)

Epitope GSDTITLPCRIKQFINMWQE

Immunogen in vitro stimulation or selection

Species (MHC) human (DRA CD4+)

References Bouhdoud et al. 2000

- CTL were studied through PBMC stimulation in vitro by gp120 pulsed autologous monocytes.
- Human CD4+ CTL clone (Een217) is an MHC class II HLA-DRA restricted CTL clone that can lyse antigen presenting HLA-DRA-transfected murine L cells - natural variants of the epitope resulted in an anergic response.
- · Low concentrations of the HXB2-derived variant (GSDTI-TLPCRIKQIINMWQK) induced T cell anergy - higher concentrations could induce proliferation and cytotoxic activity.
- (TGDIITLPCRIKQII-NRWQV), Eli (TNT-NITLQCRIKQIIKMVAG) and Z3 (CTGNITLPCRIKQIIM-NWQE) variants did not induce proliferation, cytotoxic or anergic responses.

HXB2 Location gp160 (416-424)

**Author Location** Env (413–421 SF2)

Epitope LPCRIKQII Immunogen HIV-1 infection Species (MHC) human (B\*5101)

> **Keywords** subtype comparisons, rate of progression References Tomiyama et al. 1999

- HLA-B27, -B51, and -B57 are associated with slow progression to AIDS, while HLA-B35, -B8, -B24 are associated with a rapid progression to AIDS (Nat. Med. 2:405, 1996;Lancet 22:1187, 1986; Hum Immunol 22:73, 1988; Hum Immunol 44:156, 1995)
- 15% of Japanese populations carry HLA-B51 while HLA-B27 and -B57 are detected in less than 0.3%
- Of the 172 HIV-1 peptides with HLA-B\*5101 anchor residues, 33 bound to HLA-B\*5101, seven of these peptides were reactive with CTL from 3 B\*5101 positive individuals, and six were properly processed.
- Four of the six epitopes were highly conserved among B subtype sequences, LPCRIKQII is not conserved.

HXB2 Location gp160 (416-424) Author Location gp160 (416–424 LAI) Epitope LPCRIKQII

Subtype B **Immunogen** 

Species (MHC) human (B\*5101)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*5101 epitope.

HXB2 Location gp160 (416-424)

**Author Location** gp120 (378–385)

Epitope LPCRIKQII

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B51)

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2001a

• ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** gp160 (416–424)

**Author Location** gp160 (416–429)

Epitope LPCRIKQII **Immunogen** HIV-1 infection Species (MHC) human (B51)

**Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DO7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these epitopes, but this increased over time.

HXB2 Location gp160 (416-429)

Author Location gp120 (410–429 H3DCG)

Epitope LPCRIKQFINMWQE

Immunogen HIV-1 infection

Species (MHC) human (DR4 CD4+)

References Siliciano et al. 1988

• CD4+ CTL restricted by class II HLA-DR4, targets primed by CD4 mediated uptake of gp120.

**HXB2 Location** gp160 (416–435)

Author Location gp120 (421–440 LAI)

Epitope LPCRIKQFINMWQEVGKAMY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Dadaglio et al. 1991

• Defined through blocking CTL activity, and Env deletions.

HXB2 Location gp160 (419-427) Author Location gp120 (424–432 HXB2) **Epitope** RIKQIINMW Subtype B Immunogen

Species (MHC) human (A\*3201) References Harrer et al. 1996b

• C. Brander notes that this is an A\*3201 epitope in the 1999 database.

HXB2 Location gp160 (419–427) Author Location gp120 (419–427 HXB2) Epitope RIKQIINMW Subtype B Immunogen

Species (MHC) human (A\*3201) Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is an A\*3201 epitope.

**HXB2 Location** gp160 (419–427) Author Location gp120 (419–427) Epitope RIKQIINMW? Immunogen HIV-1 infection Species (MHC) human (A29, A32) Keywords immunodominance References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immuno-
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals was A29 and responded to RIKOI-INMW, and another responder was A32 and these are thought to be presenting molecules.
- The sequence is unclear Betts calls both peptide 30 and peptide 32 gp120 419–427 and the peptide sequences are not provided.

HXB2 Location gp160 (419-427) Author Location gp120 (424-432 LAI) Epitope RIKQFINMW Subtype B Immunogen HIV-1 infection

Species (MHC) human (A32)

References Ray et al. 1998

- Autologous virus was used to detect CTL in two individuals, and in both cases strain-specific autologous CTL were found.
- The autologous epitope sequence was RIKQIINMW, MN and RF were KIKQFINMW and RIKQFVNMW respectively, and all were reactive with CTL clones.

**HXB2 Location** gp160 (419–427) Author Location gp120 (420–428) Epitope RIKQIINMW Immunogen HIV-1 infection Species (MHC) human (A32) References Ferris et al. 1999

• This epitope is processed by a TAP1/2 dependent mechanism.

**HXB2 Location** gp160 (419–427) Author Location gp120 Epitope RIKQIINMW Epitope name A32-RW10(gp120) Subtype B

Immunogen HIV-1 infection Species (MHC) human (A32)

**Donor MHC** A32, B44; A30, A32, B18, B27

Keywords HAART, ART, supervised treatment interruptions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hivweb.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples of the 15 patients, and an additional 8 responses were detected only in LN. The total magnitude of the response was similar in LN and PB, but the percentage of CD8+ T cells in the LN is lower so the number of HIV-specific cells per million CD8+ T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals. Patient B displayed the greatest response to epitope B44-AW11(p24) and also responded to A32-PW10(RT) in both PB and LN samples, while a third response against epitope A32-RW10(gp120) was only detected in the LN sample. Patient D displayed the greatest response to B27-KK10 (p24), and also responded to A30-RY11(p17), A32-PW10(RT), A30-KY11(RT), A32-RW10(gp120), and B18-YY9(Nef).

**HXB2 Location** gp160 (419–427) **Author Location** Env (424–432 BRU)

> Epitope RIKQIINMW Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (A32)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFN $\gamma$ **Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the
- number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 0/9 CRF02\_AG-infected Ivorians, and 3/9 B-infected French subjects.

HXB2 Location gp160 (421–435) Author Location gp120 (421-440 LAI) **Epitope** KQFINMWQEVGKAMY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Dadaglio et al. 1991

• Defined through blocking CTL activity, and Env deletions.

**HXB2 Location** gp160 (421–436)

Author Location gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Immunogen HIV-1 infection Species (MHC) human (A2)

**References** Clerici *et al.* 1991a

 Helper and cytotoxic T cells can be stimulated by this peptide (T1)

HXB2 Location gp160 (421–436)

Author Location gp120 (428-443 IIIB)

Epitope KQIINMWQEVGKAMYA

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Cease et al. 1987

 Helper and cytotoxic T cells can be stimulated by this peptide (T1)

**HXB2 Location** gp160 (421–436)

Author Location gp120 (428-443 IIIB)

Epitope KQIINMWQEVGKAMYA

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species (MHC)** mouse (H-2<sup>a</sup>, H-2<sup>b</sup>, H-2<sup>f</sup>)

References Shirai et al. 1992

 In a murine system multiple class I molecules can present to CTL.

**HXB2 Location** gp160 (421–436)

Author Location gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Immunogen HIV-1 exposed seronegative

Species (MHC) human

References Pinto et al. 1995

 CTL and T helper cell reactivity in healthcare workers exposed to HIV.

**HXB2 Location** gp160 (421–436)

Author Location gp120 (MN)

Epitope KQIINMWQEVGKAMYA

Immunogen HIV-1 infection

Species (MHC) chimpanzee

References Lubeck et al. 1997

• Epitope-specific CTL detected in chimpanzees immunized with

adenovirus-HIV-1 MN gp160 recombinant.

 CTL response may account for protection against subsequent HIV-1 SF2 challenge in a chimpanzee lacking neutralizing antibodies.

• Helper and cytotoxic T cells can be stimulated by this peptide (T1)

HXB2 Location gp160 (425-434)

**Author Location** Env

Epitope NMWQEVGKAM

Epitope name 1257

Subtype multiple

Immunogen HIV-1 infection

Species (MHC) human (A2)

Donor MHC A02, A30, B39; A02, A03, B44, Cw05, Cw07

Country United States.

Assay type T-cell Elispot

Keywords binding affinity, computational epitope predic-

tion

References De Groot et al. 2003

- Epitopes defined using sequence parsing and matching algorithm Conservatrix, and epitope prediction tool EpiMatrix, were shown to be conserved in a broad range of HIV-1 sequences derived from different parts of the world. 31 novel highly conserved HIV-1 epitopes were found, among which four were recognized as promiscuous epitopes and five as MHC supertypes.
- Estimated binding probability for NMWQEVGKAM: 50%

**HXB2 Location** gp160 (432–451)

Author Location gp120 (439–458 IIIB)

Epitope KAMYAPPISGQIRCSSNITG

Immunogen vaccine

Vector/Type: virus-like particle (VLP) HIV

component: CD4BS, Gag, gp120, V3

Species (MHC) macaque

References Wagner et al. 1998b

- A VLP is a non-infectious virus-like particle self-assembled from HIV Pr55 gag – macaques were immunized with VLPs bound to either gp120 or V3+CD4 linear domains Gag and Env specific CTL were stimulated in each case, and Ab response to gag and gp120 was elicited, but the gp120 neutralizing response occurred only with whole gp120, not V3+CD4 – despite the CTL and Ab response, immunized macaques were infected by intervenous challenge with SHIV chimeric challenge stock.
- CTL specific for this epitope could be found both before and after SHIV challenge.

HXB2 Location gp160 (434–443)

**Author Location** gp120 (431–440)

Epitope MYAPPIGGQI

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2K<sup>d</sup>)

References Duarte et al. 1996

 Tolerization of CTL response with continued administration of soluble peptide.

**HXB2 Location** gp160 (435–443)

**Author Location** 

Epitope YAPPISGQI

Immunogen SHIV infection

Species (MHC) macaque (Mamu-A\*01)

References Egan et al. 1999

SHIV-infected rhesus macaques have high frequencies of response to the SIVmac epitope gag p11C,C-M (CTPYDINQM) but only a fraction of A\*01 monkeys tested have responses to SIVmac pol epitope STPPLVRLV and HIV-1 env epitope YAPPISGQI.

**HXB2 Location** gp160 (435–443)

Author Location gp41 (89.6)

Epitope YAPPISGQI

Epitope name p41A

Immunogen SHIV infection, vaccine

Vector/Type: DNA, modified vaccinia Ankara (MVA) Strain: B clade 89.6, B clade HXBc2 HIV component: Env, Gag Adju-

vant: IL-2/Ig

Species (MHC) macaque (Mamu-A\*01)

**Keywords** immunodominance **References** Barouch *et al.* 2001a

- Mamu-A\*01+ rhesus monkeys infected with SHIV-89.6 and SHIV-HXBc2 make immunodominant responses to SIV Gag p11C epitope (CTPYDINQM) and a subdominant response to HIV-1 Env p41A epitope (YAPPISGQI)
- The binding affinities are the same for the two Mamu A\*01 epitopes, so that is not what dictates the dominance.
- Monkeys vaccinated with MVA vectors carrying SIV gag/pol and HIV-1 env showed the same p11C epitope dominance and p41A epitope subdominance, but co-dominance was observed and the response to p41A increased when DNA vaccination was done using the SIV and HIV genes under CMV promotor control with IL2-IG adjuvant.

**HXB2 Location** gp160 (435–443)

**Author Location** Env

**Epitope** YAPPISGQI **Immunogen** vaccine

Vector/Type: DNA Strain: B clade 89.6P,

SIV HIV component: Env, Gag

Species (MHC) macaque (Mamu-A\*01)

Assay type Flow cytometric T-cell cytokine assay

**Keywords** vaccine-specific epitope characteristics, rate of progression, kinetics, memory cells, char-

acterizing CD8+ T cells

References Davenport et al. 2004

• Activation and expansion of antigen-specific CD8+ T-cells shows a delay following infection that allows early viral replication. Until day 10, the kinetics of CD8+ T-cell expansion was the same in vaccinated and control macaques. An increase in virus-specific CD8+ T-cell numbers around day 10 in vaccinated macaques coincides with a slowing in viral replication. This indicates that while cytotoxic T-lyphocyte-inducing vaccines may have a long-term benefit in controling viral replication and preventing disease progression, they cannot prevent infection.

**HXB2 Location** gp160 (435–443)

**Author Location** Env (89.6)

Epitope YAPPISGQI

Epitope name p41A

Immunogen vaccine

Vector/Type: DNA Strain: B clade 89.6, SIV HIV component: Env, Gag Adjuvant:

IL-2/Ig **Species (MHC)** macaque

References Barouch et al. 2000; Shen & Siliciano 2000

 Different HIV strains were used for different regions: SIVmac239 Gag and HIV-1 89.6P Env

- Monkeys that received the DNA vaccines augmented with IL-2/Ig were infected when challenged with pathogenic SHIV-89.6P, but had potent CTL responses, stable CD4+ T-cell counts, preserved virus-specific CD4+ T-cell responses, low to undetectable viral loads, and no evidence of disease or mortality by day 140 after challenge – monkeys that got a sham vaccine had high viral load, progressed to disease, and were half were dead by day 140.
- IL2/Ig consisting of interleukin-2 (IL-2) for immune stimulation, and the Fc portion of immunoglobulin G (IgG) for stability, was delivered either as protein or as DNA both enhance the CTL response to vaccination, DNA IL2/Ig giving the most intense response.
- Responses to a dominant Mamu A\*01 gag epitope SIV Gag p11C (CTPYDINQM) and a subdominant epitope HIV-1 Env p41A (YAPPISGQI) were tracked and had good durability prior to challenge, and the higher the prechallenge peak p11C CTL response, the lower the post-challenge viral load.
- No NAb responses were detected in the vaccinated monkeys prior to challenge, and comparable peak NAb titers developed in vaccinated monkeys and control monkeys with preserved CD4+ T-cells.
- Shen et al. 2000 is an accompanying commentary.

**HXB2 Location** gp160 (435–443)

**Author Location** Env (89.6)

Epitope YAPPISGQI

Epitope name p41A

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade 89.6, SIV HIV component: Env, Gag-Pol Adju-

*vant:* IL-2/Ig

Species (MHC) macaque

Keywords immunodominance

References Barouch et al. 2001b

- Different HIV strains were used for different regions: SIV-mac239 Gag/Pol and HIV-1 89.6P Env
- Four monkeys were vaccinated with a modified vaccinia Ankara (MVA) vaccine that elicited strong CTL to the immunodominant SIV gag epitope in 4/4 animals, and 1/4 made a response to the HIV Env epitope YAPPISGQI, as determined by tetramer staining and chromium release assays.
- The animals were infected when challenged with pathogenic SHIV-89.6P, but had potent CTL responses, secondary NAb responses upon challenge, partial preservation of CD4+ T-cell counts, lower viral loads, and no evidence of disease or mortality by day 168 after challenge monkeys that got a sham vaccine had high viral load, progressed to disease, and 2/4 were dead by day 168.

HXB2 Location gp160 (444–453)

Author Location Env

Epitope RCSSNITGLL

Immunogen

Species (MHC) human (B56)

References De Groot et al. 2001

The program Epimatrix was used in conjunction with the program Conservatrix to identify conservered regions of HIV that might serve as epitopes.

- A subset of the potential epitopes was identified that could bind to the appropriate HLA-allele, and 15 of the predicted B7 superfamily (HLA B7, B8, and B58) epitopes were identified that could stimulate IFNγ production in an ELISPOT assay.
- RCSSNITGLL was newly defined as an epitope in this study, and was shown to stimulate an ELISPOT response, despite not detectably binding to HLA-B7.

**HXB2 Location** gp160 (466–475)

**Author Location** Env (464–473)

Epitope EVFRPGGGDM

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*2601)

Country Japan.

**Assay type** Chromium-release assay, Flow cytometric T-cell cytokine assay, Other, HLA binding

Keywords immunodominance, optimal epitope

References Satoh et al. 2005

- Reverse immunogenetics was used to identify HIV-1 epitopes presented by HLA-A\*2601. 110 peptides were predicted to bind to HLA-A\*2601. 24 of these were demonstrated to bind through a HLA-A\*2601 stabilization assay. Four of these, including this one, were shown to be epitopes endogenously presented by this allele, that can induce peptide-specific CD8 T-cells.HLA-A\*2601 is common in Asia.
- This epitope was recognized in only 1/7 HLA-A\*2601 HIV infected individuals.

HXB2 Location gp160 (486–494)

Author Location gp160 (485–493 SUMA)

**Epitope** YKVVKIEPL **Epitope name** GP160 YL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*1103, A\*2402, B\*1402, B\*1501,

Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, acute/early infection, characterizing CD8+ T cells

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient SUMA maintained low viral loads and stable CD4 T cell counts through seven years of follow up. In contrast to more rapid progressors, WEAU and BORI, SUMA a broad response to 24 epitopes, with little immunodominance. Two peptides were somewhat more intensely recognized in acute infection, but this response leveled out early on.

• Only four epitopes were found to acquire escape muations in SUMA over time, and this was one of the 20 that remained invariant. A low level response was detected at acute infection that persisted through early infection.

**HXB2 Location** gp160 (489–508)

Author Location gp120 (494-513 BRU)

Epitope VKIEPLGVAPTKAKRRVVQR

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Dadaglio et al. 1991

• Defined through blocking CTL activity, and Env deletions.

**HXB2 Location** gp160 (489–508)

**Author Location** Env (501–)

Epitope VKIEPLGVAPTKAKRRVVQR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A\*0202, A\*0301, B\*0702, B\*1516

Country United States.

Keywords escape, acute/early infection

References Bernardin et al. 2005

- Full HIV genomes from nine individuals were analyzed for mutations prior to seroconversion and 7 to 28 days later. It was found that the influence of the host's HLA type was reflected within weeks of infection; a statistically significant number of early nonsynonymous mutations were observed within previously reported CTL epitopes.
- K to R mutation was observed in position 16.

HXB2 Location gp160 (489–508)

Author Location Env (496-506 BH10, LAI)

Epitope VKIEPLGVAPTKAKRRVVQR

Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is VAPTKAKR-RVV) has similarity with the mast/stem cell growth factor receptor precursor fragment VVPTKADKRRSV.

HXB2 Location gp160 (489–508)

Author Location Env (497–512 BH10, LAI)

Epitope VKIEPLGVAPTKAKRRVVQR

Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is APTKAKR-RVVQREKRA) has similarity with the human interferonrelated IFRD2 (PC4-B) protein fragment ARTKARSRVRD-KRA.

**HXB2 Location** gp160 (511–519)

Author Location gp160 (511–519)

Epitope YRLGVGALI

Epitope name YI9

Immunogen

Species (MHC) human (Cw\*18)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a Cw18 epitope.

**HXB2 Location** gp160 (511–519)

Author Location (C consensus)

Epitope YRLGVGALI

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*1801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- YRLGVGALI is an optimal epitope.

**HXB2 Location** gp160 (511–526)

**Author Location** (C consensus)

Epitope RAVGIGAVFLGFLGAA

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** gp160 (519–543)

Author Location gp41 (519–543)

Epitope FLGFLGAAGSTMGAASLTLTVQARC

Immunogen HIV-1 infection

Species (MHC) human (Cw7)

References Nehete et al. 1998a

- Three long-term non-progressors and one asymptomatic HIV+ individual were studied and found to have HLA class I Crestricted CD8+ Env-specific CTLs – Cw7 specific CTL were found against three peptides, including this one.
- HLA-C antigens are expressed on lymphoid cells to a lesser extent, 10% of either HLA-A or HLA-B.
- HLA-C confers protection against lysis by natural killer cells and by non-MHC-restricted effector T cells and Cw7 directly governs this resistance to lysis – the authors hypothesize that pathogens that inhibit antigen expression and class I expression may particularly down regulate Cw7, thus triggering non-MHC restricted killing.

HXB2 Location gp160 (529-537)

Author Location Env (529-)

Epitope TMGAASITL

Epitope name Env529

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: gp160 Adjuvant: Incomplete Freund's Ad-

juvant (IFA)

Species (MHC) human (A2)

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was an intermediate A2 binder that did not induce CTL and CD8+ T-cell IFN gamma responses in mice.
   Responses were detected in 5/17 HIV+ HLA-A2 subjects.

**HXB2 Location** gp160 (552–571)

**Author Location** Env (552–571)

Epitope QSNLLRAIEAQQHMLQLTVW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** gp160 (557–564)

**Author Location** (C consensus)

Epitope RAIEAQQM

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- RAIEAQQM is an optimal epitope.

**HXB2 Location** gp160 (557–565)

Author Location gp41 (557–665)

Epitope RAIEAQQWQ

Epitope name E3

Immunogen HIV-1 infection Species (MHC) human (B\*5101) Keywords HAART, ART, escape References Samri et al. 2000

• The epitope was recognized by patient 246#1 in a study of the effects of therapy escape mutations on CTL recognition.

HXB2 Location gp160 (557–565) **Author Location** gp41 (46–54 HIV-MN)

Epitope RAIEAQQHL

Epitope name RL9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*5101)

**Donor MHC** A\*0201, A\*2501, B\*5101, B\*1801,

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords optimal epitope References Liu et al. 2006

• T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.

HXB2 Location gp160 (557-565)

**Author Location** gp160 (557–565)

Epitope RAIEAQQHL Immunogen HIV-1 infection Species (MHC) human (B15, B51)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4, DR8, DO7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

HXB2 Location gp160 (557-565)

Author Location gp41 (557–565 IIIB)

Epitope RAIEAQQHL Immunogen HIV-1 infection Species (MHC) human (B51) References Sipsas et al. 1997 • HIV IIIB proteins were used to define the range of CTL epitopes recognized by 3 lab workers accidentally infected with HIV-1 IIIB.

- · KAIEAQQHL, a variant found in HIV-1 NY5CG, was also recognized.
- RAIEAOOHM, a variant found in HIV-1 JRCSF, was also recognized.
- RAIDAOOHL, a variant found in HIV-1 ETR, was also recognized.
- RAIKAQQHL, a variant found in HIV-1 CDC42, was also recognized.

**HXB2 Location** gp160 (557–565)

Author Location gp41 (557–565)

Epitope RAIEAQQHL

Immunogen HIV-1 infection

Species (MHC) human (B51)

References Ferris et al. 1999

• This epitope can be processed by a TAP1/2 dependent mecha-

**HXB2 Location** gp160 (557–565)

Author Location gp41 (557–565)

Epitope RAIEAQQWQ

Epitope name RAI

Immunogen HIV-1 infection

Species (MHC) human (B51)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- None of the 8 study subjects recognized this epitope but none were HLA B51+

HXB2 Location gp160 (557–565)

Author Location gp41 (47–55)

Epitope RAIEAQQHL

Immunogen HIV-1 infection

Species (MHC) human (B51)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** gp160 (557–565)

**Author Location** gp41 (557–565 LAI)

Epitope RAIEAQQHL

Epitope name E3

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B51)

Keywords HAART, ART

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+PBL but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

**HXB2 Location** gp160 (557–565)

Author Location gp41

Epitope RAIEAQQHL

Subtype B, C

Immunogen HIV-1 infection

**Species (MHC)** human (B63, B57, B58)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords cross-presentation by different HLA, optimal

epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This is a putative HLA-B63/57/58 epitope containing the B58 supertype binding motif. Significantly more often recognized by B63+ and B57/58+ subjects than by negative subjects.

HXB2 Location gp160 (557–565)

Author Location Env (gp160) (557-565)

Epitope RAIEAQQHL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0304)

**Keywords** subtype comparisons

**References** Currier *et al.* 2002a

- Cross-reactive responses were found in PBMC isolated from individuals infected with either B or CRF01\_AE clade viruses, as determined by Elispot assays of target cells expressing recombinant vaccinia viruses expressing HIV-1 gag, env, nef and pol from many clades.
- CTL from subject US101, infected with a clade B virus, displayed broad cross-reactivity to HIV-1 clade A, B, C, D, CRF01\_AE, F G, recognized this epitope. Clade B and C had a L->M change in the C-term position that was tolerated. The H clade Env was not cross-reactive, and had the sequence RAIAARQHM.

**HXB2 Location** gp160 (557–565) **Author Location** gp41 (46–54)

Epitope RAIEAQQHL

Immunogen

Species (MHC) human (Cw\*0304)

**Keywords** optimal epitope

References Frahm et al. 2007

**HXB2 Location** gp160 (557–565)

Author Location (C consensus)

Epitope RAIEAQQHM

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords cross-presentation by different HLA, charac-

terizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** gp160 (557–565)

Author Location gp41 (46-54)

Epitope RAIEAQQHL

Immunogen

Species (MHC) human (Cw\*15)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** gp160 (557–565)

**Author Location** 

Epitope RAIEAQQHM

Epitope name RM9

Immunogen

Species (MHC) human (Cw8)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a Cw08 epitope.

HXB2 Location gp160 (557–565)

Author Location gp41 (557–565 IIIB)

Epitope RAIEAQQHL

Immunogen HIV-1 infection

Species (MHC) human

Keywords responses in children, mother-to-infant trans-

mission

References Wilson et al. 1996

- Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.
- RAIDAQQHL and RVIEAQQHL, naturally occurring variants, were found in mother and are recognized.

**HXB2 Location** gp160 (557–565) **Author Location** gp41 (557–565)

**Epitope** RAIEAQQHL **Immunogen** HIV-1 infection

Species (MHC) human

**Keywords** immunodominance **References** Betts *et al.* 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INFγ responses to other epitopes.
- 1/11 of the A2+ individuals was HLA A\*0201, A32, B60, B78, and responded to RAIEAQQHL, previously noted to be B51.

**HXB2 Location** gp160 (557–565) **Author Location** gp41 (557–565 IIIB)

Epitope RAIEAQQHL Immunogen HIV-1 infection

Species (MHC) human

Keywords mother-to-infant transmission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- This epitope was invariant in both the mother and her infant.

HXB2 Location gp160 (557–565)

Author Location Env (555–567 BH10, LAI)

Epitope RAIEAQQHL Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is LL-RAIEAQQHLL) has similarity with human MHC class II regulatory factor RFX1 fragment LLRLMEDQQHMA.

**HXB2 Location** gp160 (557–565)

**Author Location** gp160

Epitope RAIEAQQHL Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: vaccinia Strain: A clade, B clade, D clade NDK, C clade consensus HIV component: Env

Species (MHC) human

**Donor MHC** A\*3201, A\*3601, B\*5301, B\*8101, Cw\*0401, Cw\*0804

Country Kenya.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Other

**Keywords** subtype comparisons, variant cross-

recognition or cross-neutralization

References McKinnon et al. 2005

- Interclade cross-reactivity of clades A, B, C, and D was tested using recombinant vaccinia-based IFN-gamma Elispot assay. 47/74 women had a positive Elispot response to at least one clade, and cross-clade responses were frequent. Clade A responses were most frequent, as expected, as clade A dominates the Kenyan epidemic (85% of the 47 women responded to clade A Env, 74% to B, 62% to C, and 43% to D). Cross-clade CD8 T-cell responses were common and directed at conserved epitopes.
- There was a greater magnitude of response to A clade peptides in individuals who responded to more than one clade; a 2-fold higher response was observed in clade A in 36% (9/25) of these individuals, and the response to A peptides was never lower. VTEEFNMWK responses were detected in 2 women that had Env responses to all 4 clades, and clade A gave the highest responses; a VnEEFNMWK variant was in clade B and D, and the clade C Env carried VnEEFNMWe. One woman also reacted with RAIEAQQHL, the other with KNCSFNMTT. RAIEAQQHL was identical in clades A, B, and D, while C carried RAIEAQQHm.

HXB2 Location gp160 (557-566)

**Author Location** Env (557–566)

Epitope RAIEAQQHML

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*40, A\*69)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** gp160 (565–573)

**Author Location** Env (565–)

Epitope LLQLTVWGI

Epitope name Env565

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Env Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (MHC) human, transgenic mouse (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced CD8+ T-cell IFN gamma responses in mice. Responses to the peptide were not detected in 17 HIV+ HLA-A2 subjects.

HXB2 Location gp160 (565–573) Author Location Env (731–739) Epitope LLQLTVWGI Immunogen HIV-1 infection

**Species (MHC)** human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind four of the five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802).

**HXB2 Location** gp160 (570–589) **Author Location** gp41 (571–590 LAI)

Epitope VWGIKQLQARILAVERYLKD

Subtype B Immunogen vaccine

Vector/Type: vaccinia prime with gp160 boost Strain: B clade LAI HIV component: gp160

Species (MHC) human (DR-1 CD4+ CTL)

References Kent et al. 1997a

- VWGIKQLQARILAVERYLKD, present in HIV-1 LAI, was the immunizing strain.
- VWGIKQLQARVLAVERYLKD, present in HIV-1 MN, was also recognized.
- VWGIKQPQARVLAVERYLRD was the form carried by the autologous strain that infected the vaccinee.
- Lysis of the target cells by CD4+ CTL was inhibited with the addition of the peptide representing the autologous strain.
- The infecting virus epitope also antagonized the proliferative functions of the CD4+ CTL clone.
- The behavior of the autologous strain presents a possible mechanism for vaccine failure since the infecting virus not only escapes CTL activity, but inhibits the ability of CTL to recognize other variants.

HXB2 Location gp160 (572–590) Author Location gp41 (572–590 BRU) Epitope GIKQLQARILAVERYLKDQ

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU HIV component: gp160

Species (MHC) human (DPw4.2)

References Hammond et al. 1991

CD4. CTI

• CD4+ CTL.

**HXB2 Location** gp160 (575–599) **Author Location** gp41 (575–599 IIIB)

Epitope QLQARILAVERYLKDQQLLGIWGCS

Immunogen HIV-1 infection Species (MHC) human (B14) References Jassoy *et al.* 1992

• Epitope recognized by CTL clone derived from CSF.

HXB2 Location gp160 (577–587) Author Location (C consensus) Epitope QTRVLAIERYL

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*5802) Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- · QTRVLAIERYL is an optimal epitope.

**HXB2 Location** gp160 (583–592)

Author Location gp41 (583–592 PV22)

Epitope VERYLKDQQL
Immunogen HIV-1 infection

Species (MHC) human (B14)

References Jassoy et al. 1993

• HIV-1 specific CTLs release  $\gamma$ -IFN, and  $\alpha$ - and  $\beta$ -TNF.

**HXB2 Location** gp160 (584–592)

Author Location gp41 (584–592 HXB2)

Epitope ERYLKDQQL

Epitope name E4

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (A32, B14)

Keywords HAART, ART References Mollet *et al.* 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+PBL but with continued viral suppression, HIV-specific responses diminished.

 Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

HXB2 Location gp160 (584-592)

Author Location gp41

Epitope ERYLRDQQL

Immunogen HIV-1 infection

Species (MHC) human (B\*14)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

**HXB2 Location** gp160 (584–592)

**Author Location** (C consensus)

Epitope ERYLKDQQL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*14)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location gp160 (584–592)

Author Location (C consensus)

Epitope ERYLKDQQL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*1401)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- ERYLKDQQL is an optimal epitope.

HXB2 Location gp160 (584–592)

Author Location gp41 (584-592 PV22)

Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human (B\*1402)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1402 epitope.

**HXB2 Location** gp160 (584–592)

Author Location gp160 (598–597 BORI, SUMA)

Epitope ERYLKDQQL

Epitope name gp160 EL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1402)

**Donor MHC** A\*2902, B\*1402, Cw\*0802; A\*1103,

A\*2402, B\*1402, B\*1501, Cw\*0802

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

**Keywords** dynamics, immunodominance, escape,

acute/early infection, characterizing CD8+ T

cells, reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient BORI had high viral loads and rapid CD4 decline. BORI mounted 8 detected responses. BORI did not control viral replication well, and escape mutations occurred early and in most epitopes, 6/7 that were precisely identified.
- The patient SUMA maintained low viral loads and stable CD4 T cell counts through seven years of follow up. In contrast to more rapid progressors, WEAU and BORI, SUMA a broad response to 24 epitopes, with little immunodominance. Two peptides were somewhat more intensely recognized in acute infection, but this response leveled out early on.
- Eleven variants in the ERYLKDQQL epitope were found in the patient BORI. ERYLKeQQp came up first at day 17 from onset of symptoms, but wasn't tested for escape properties. ERYLrDQQL came up next, by day 31, but didn't confer escape in a Cr release assay. By day 218, three variants were found, all of which gave a diminished response: ERYLtDQQL, ERYLqDQQL, and ERYLsDQQL. By day 556 a complex mixture was present, also including the ERYLmDQQL variant

that gave a further reduction in the response, and many double mutants: ERYLmDQrL, ERYLmDrQL, ERYLmDQlL, ERYLtDOrL and ERYrtDOrL.

• In SUMA, the only variation found in the 24 epitopes was in three overlapping epitopes in Tat, and in this gp160 epitope; variation accumulated early in infection in the Tat epitopes, but this epitope was stable until a sample 736 days post-infection, when only the ERYLqDQQL variant was detected. This variant was not tested with CTL from SUMA, but gave a diminished response in BORI.

**HXB2 Location** gp160 (584–592)

Author Location gp41

Epitope ERYLKDQQL
Immunogen HIV-1 infection
Species (MHC) human (B14)

References Wagner et al. 1998a

• CTL specific for HIV epitopes were used to show that the mediators of both the cytolytic (granzyme A was used as the marker) and non-cytolytic (HIV-1 inhibitory chemokines MIP-1  $\alpha$  and RANTES were used as markers) anti-viral responses are localized within the CTL's cytotoxic granules.

HXB2 Location gp160 (584–592) Author Location gp41 (584–592) Epitope ERYLKDQQL

Immunogen HIV-1 infection Species (MHC) human (B14) Keywords HAART, ART

References Kalams et al. 1999b

- Two patients were followed before and after HAART reduced plasma HIV-1 RNA levels resulted in a decline in HIV in vivo activated specific CTL such that by day 260 CTL activities were undetectable.
- ERYLKDQQL was the dominant response in one of the individuals, SLYNTVATL subdominant.
- Sporadic breakthrough in viremia resulted in increases in CTLp.
- Peptide-tetramer staining demonstrated that declining levels of *in vivo*-activated CTL were associated with a decrease in expression of CD38.
- Memory CTL frequency directed against Vac-Gag, Vac-RT, Vac-Env, and Vac-Nef initially increased with HAART and then decreased with the decline of the viral load.

**HXB2 Location** gp160 (584–592)

Author Location gp41 (591–599 SF2)

**Epitope** ERYLKDQQL **Immunogen** HIV-1 infection

Species (MHC) human (B14)

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-A3, -A32, -B7, -B14.

**HXB2 Location** gp160 (584–592) **Author Location** gp41 (591–599 SF2) Epitope ERYLKDQQL
Immunogen HIV-1 infection
Species (MHC) human (B14)
Keywords subtype comparisons

References Cao et al. 1997a

- The consensus sequence for clades B, C, and D is ERYLKDQQL.
   The consensus sequence for clade A is ERYLBDOOL and it is
- The consensus sequence for clade A is ERYLRDQQL and it is equally reactive.
- The consensus sequence for clade E is ERYLKDQKF and it is not reactive.

**HXB2 Location** gp160 (584–592)

**Author Location** gp41

Epitope ERYLKDQQL

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B14)

**Keywords** subtype comparisons, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades such cross-reactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A and D subtype consensus are identical to the B clade epitope, ERyLkDQQL.

HXB2 Location gp160 (584–592)

Author Location gp41 (584–592)

Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Sipsas et al. 1997

 HIV IIIB proteins were used to define the range of CTL epitopes recognized by 3 lab workers accidentally infected with HIV-1 IIIB.

HXB2 Location gp160 (584–592)

**Author Location** gp41 (584–592)

Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Yang et al. 1996

- CD4+ cell lines acutely infected with HIV were studied to determine their susceptibility to lysis by CTL.
- Clones specific for RT lysed HIV-1 infected cells at lower levels than Env or Gag specific clones.
- The distinction was thought to be due to lower expression of RT relative to Env and Gag.
- CTL can lyse infected cells early after infection, possibly prior to viral production.

HXB2 Location gp160 (584–592)

**Author Location** gp41 (584–592)

Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human (B14)

Assay type CTL suppression of replication

References Yang et al. 1997a

- comparable to those found in vivo.
- MIP-1 $\beta$ , RANTES, after antigen-specific activation.
- CTL suppress HIV replication more efficiently in HLA- 3/5 subjects showed no variation in viral sequence, 2/5 matched cells.

HXB2 Location gp160 (584-592) Author Location gp41 (584-592 PV22) Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human (B14) References Johnson et al. 1992

• Two overlapping CTL epitopes were mapped with different HLA restriction (also see YLKDQQLL HLA-B8)

**HXB2 Location** gp160 (584–592)

Author Location gp41 (584–592 PV22)

Epitope ERYLKDQQL Immunogen HIV-1 infection Species (MHC) human (B14)

References Jassoy et al. 1993

• HIV-1 specific CTLs release  $\gamma$ -IFN, and  $\alpha$ - and  $\beta$ -TNF.

HXB2 Location gp160 (584–592)

Author Location gp41 (584–592 HXB2)

Epitope ERYLKDQQL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B14)

References Kalams et al. 1994; Kalams et al. 1996

- Longitudinal study of T cell receptor usage in a single individ- CTL responses in seronegative highly HIV-exposed African
- Persistence of oligoclonal response to this epitope for over 5 years.

HXB2 Location gp160 (584-592)

Author Location gp41 (584–592)

Epitope ERYLKDQQL

Immunogen peptide-HLA interaction

Species (MHC) human (B14)

References DiBrino et al. 1994a

· Epitope studied in the context of HLA-B14 binding.

HXB2 Location gp160 (584–592)

Author Location gp41 (584–592)

Epitope ERYLKDQQL Immunogen HIV-1 infection

Species (MHC) human (B14)

References Hammond et al. 1995

• This peptide can be processed for HLA-B14 presentation in a TAP-1/2 independent pathway.

HXB2 Location gp160 (584-592)

Author Location gp41 (584–592)

Epitope ERYLKDQQL Immunogen HIV-1 infection Species (MHC) human (B14)

References Kalams et al. 1996

• CTL response to this epitope was studied in 5 HLA-B14 positive persons.

- CTL inhibit HIV-1 replication at effector cell concentrations CTL responses were detected in all five, and CTL clones were isolated from 4/5.
- CTL produced HIV-1-suppressive soluble factors MIP- $1\alpha$ , A diverse repertoire of TCRs recognized this epitope, with similar fine specificities.
  - had a dominant variant that resulted in poor recognition, ERYLQDQQL.
  - A minor CTL response specific for the ERYLODOOL could be detected by two individuals, but the major CTL response was to the ERYLKDQQL form even when it was the minority form.
  - Some single amino acid substitutions were well tolerated by most of the CTL clones tested, but others, particularly in the center three amino acid positions, abrogated peptide stimulatory activity.

HXB2 Location gp160 (584–592)

**Author Location** gp120 (584–592)

Epitope ERYLKDQQL Immunogen HIV-1 infection Species (MHC) human (B14)

References Ferris et al. 1999; Hammond et al. 1995

· This epitope is processed by both TAP1/2 dependent and independent mechanisms.

HXB2 Location gp160 (584–592)

Author Location gp41

Epitope ERYLKDQQL

Immunogen

Species (MHC) human (B14)

References Rowland-Jones et al. 1999

- female sex workers in Gambia and Nairobi were studied these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive, and the B35 allele seems to be protective.
- HIV-2 sequence: EKYLODOAR no cross-reactivity Johnson et al. [1992]

HXB2 Location gp160 (584-592)

Author Location gp41 (SF2)

Epitope ERYLKDQQL

Epitope name EL9

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords acute/early infection

References Goulder et al. 2001a

• Data from patient AC13 suggest a role for this epitope in initial control of viremia in acute infection, as it is one of several subdominant CTL epitopes recognized during the initial decline

in viremia.

• A CTL response to SL9, SLYNTVATL, was not evident until 18 months post-presentation.

• Recognized by two A\*0201-positive chronically infected subjects.

**HXB2 Location** gp160 (584–592)

Author Location gp41 (584–592)

Epitope ERYLKDQQL

Epitope name 588K

Immunogen HIV-1 infection Species (MHC) human (B14)

Keywords HAART, ART, TCR usage

References Islam et al. 2001

- Transcript frequencies of four CTL clones from patient 115, with a chronic and stable HIV-1 infection, were tracked in a longitudinal study of samples collected 6-11 years post infection: clone M21 and E15 recognize ERYLKDQQL, and clone D87 recognizes variant ERYLQDQQL, and clone p175b recognizes the A2 epitope SLYNTVATL.
- CTL clone M21 uses the V $\beta$  4, CDR3 VKDGA, J $\beta$  1.2 TCR beta gene, and clone E15 uses the V $\beta$  4, CDR3 VEDWGGAS J $\beta$  2.1 TCR beta gene, and D87 uses V $\beta$ 8, ALNRVD, J $\beta$ 2.1.
- Responses were stable even through HAART with undetectable viral loads but frequencies varied over time by 100-fold, ranging from 0.012% of the total population for SLYNTVATL at its lowest point to 3.78% for M21, with the relative frequencies of clones shifting over time.

**HXB2 Location** gp160 (584–592)

Author Location gp41 (589–597 SF2)

Epitope ERYLKDQQL Immunogen HIV-1 infection Species (MHC) human (B14)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B14+ individuals that had a CTL response to this epitope broken down by group: 3/3 group 1, 1/2 group 2, and 0/0 group 3.

**HXB2 Location** gp160 (584–592)

**Author Location** gp41 (589–597)

 ${\bf Epitope} \ {\tt ERYLRDQQL}$ 

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B14)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** gp160 (584–592) **Author Location** gp41 (JRCSF)

Epitope ERYLKDQQL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Severino et al. 2000

- Primary HLA-B14+ CD4+ CD3+ and HLA-mismatched lymphocytes from uninfected donors were infected with JRCSF after isolation then cocultured with the B14-restricted CTL clone 15160/D75 specific for ERYLKDQQL, and viral inhibition was MHC-restricted.
- Primary monocytes and monocyte-derived DC were generated from the same donors, replication of HIV-1 in these cell types was less efficient than in lymphocytes and could also be inhibited by MHC-restricted CTL.
- DC-lymphocyte cluster cultures allowed vigorous viral replication and MHC-restricted CTL viral inhibition was blunted or lost depending on the ratio of DC to CD4+ lymphocyte in the culture.

**HXB2 Location** gp160 (584–592)

Author Location gp41 (SF2)

Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human (B14)

References Altfeld et al. 2000

 This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.

**HXB2 Location** gp160 (584–592)

**Author Location** Env (589–597)

Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Keywords** early-expressed proteins, kinetics

References Guillon et al. 2002b

 An early-expressed Nef protein was modified to contain Env and Pol epitopes to enable the study the effect of expression kinetics on CTL mediated suppression of replication. The "EpiNef" construct was inserted into a recombinant vaccinia virus which was used to infect a target cell line; the target cells were lysed by CTL clones specific for the Env and Pol epitopes indicating that they were properly processed.

**HXB2 Location** gp160 (584–592)

Author Location gp41 (584–592)

Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human (B14)

Keywords class I down-regulation by Nef

**References** Yang et al. 2002

• Nef down-modulates class I protein expression, and this study demonstrates directly that Nef-deleted HIV-1 NL-43 can be more effectively killed *in vitro* than NL-43 with an intact Nef. The effect was shown to be specific for class I presentation of epitopes, and unlike Nef, deleting Vpr did not alter CTL susceptibility of NL43 infected cells. The CTL clone 15160D75, specific for the class I B14 presented epitope ERYLKDQQL, was one of four used in this study.

HXB2 Location gp160 (584–592)

**Author Location** gp41

Epitope ERYLKDQQL

**Epitope name** B14-EL9(gp41)

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Donor MHC** A32, B7, B14

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals. Patient A displayed the greatest response to epitope B14-EL9(gp41), a strong response to B7-TL9(p24), and responses to B7-TM9(Nef) and A32-PW10(RT).

**HXB2 Location** gp160 (584–592)

Author Location gp41

Epitope ERYLKDQQL Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

Species (MHC) human (B14)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

• The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].

 Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

**HXB2 Location** gp160 (584–592)

**Author Location** gp41 (73–81)

Epitope ERYLKDQQL

Epitope name Env EL9

Subtype B

**Immunogen** HIV-1 infection **Species** (MHC) human (B14)

Assay type Chromium-release assay

**Keywords** binding affinity, TCR usage, characterizing CD8+ T cells

References Yang et al. 2003b

- Killing efficiency of CTLs and their ability to suppress viral replication is shown to depend more on epitope specificity than to antigenic avidity. Different clones recognizing the same epitope had similar killing efficiency despite their variation in avidity. Nef specific CTL clones tended to be most inhibitory, followed by Gag, then by RT specific clones, regardless of avidity.
- 4/14 CTL T-cell clones tested were specific for Env EL9. Under conditions of excess peptide (100ug/ml), there was no difference in their lytic potential; all possessed similar effector capacity. Avidity was measured as the sensitizing dose of peptide required for 50% of maximal killing (SD50), which varied from 20 pg/ml to 100 ng/ml, over four orders of magnitude for all 14 epitopes. The SD50 range for Env EL9 was 5,000 60,000 pg/ml.

HXB2 Location gp160 (584–592)

Author Location (B consensus)

Epitope ERYLKDQQL

Epitope name EL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

Donor MHC A28, A29, B14, B44, Cw8; A25, A32, B08,

B14, Cw7, Cw8

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

Using a flow-cytometric cytotoxicity assay based on caspase-3
activation in dying target cells, it was shown that the subset of
HIV-1-specific CD8+ T cells secreting both IFN-gamma and
TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
perforin expression was found.

B14.

HXB2 Location gp160 (584–592) **Author Location** Env (584–592) Epitope ERYLKDQQL

Epitope name EL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B14)

**Donor MHC** A\*02, A\*68, B\*14, B\*52, Cw\*08, Cw\*12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords escape, characterizing CD8+ T cells, optimal epitope

References Koibuchi et al. 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- The variant ERYLqDQQL was the only form of the epitope detected over a 6-year period in this person. Elispot reactions were reduced to the autologous form relative to the B clade consensus form, ERYLKDQQL.

HXB2 Location gp160 (584-592)

Author Location gp41

Epitope ERYLKDQQL

Epitope name EL9

Immunogen

Species (MHC) (B14)

Keywords review, immunodominance. escape, acute/early infection, early-expressed proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

• This review discusses the importance of 3 factors that impact the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides listed as immunodominant in acute HIV-1 infection.

HXB2 Location gp160 (584-592)

Author Location gp41 (subtype B)

Epitope ERYLKDQQL

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B14, B\*1402)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi - these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.

- 2/9 individuals recognized this epitope, presented by HLA- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found - B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
  - This epitope is conserved among B and D clade viruses.
  - The Clade A version of the epitope is ERYLRDQQL.

HXB2 Location gp160 (584-592)

Author Location gp41 (584–592)

Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human

References Price et al. 1995

• Study of cytokines released by HIV-1 specific activated CTL.

**HXB2 Location** gp160 (584–592)

**Author Location** gp41 (584–592)

Epitope ERYLKDQQL

Immunogen HIV-1 infection

Species (MHC) human

References Borrow et al. 1994

- Three out of five patients with HIV-1 symptomatic infection controlled their viral infection well and mounted an early, strong HIV-1 specific MHC restricted CTL response.
- One of the three, study subject BORI, specifically recognized this peptide.

HXB2 Location gp160 (584-592)

Author Location gp160

Epitope ERYLRDQQL

Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: vaccinia Strain: A clade, B clade, D clade NDK, C clade consensus HIV

component: Env

Species (MHC) human

Donor MHC A\*0202, A\*7401, B\*5802, B\*1503, Cw\*0202, Cw\*0602; A\*0201, A\*3009,

B\*5802, B\*4501, Cw\*0202, Cw\*1601

Country Kenya.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Other

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References McKinnon et al. 2005

- Interclade cross-reactivity of clades A, B, C, and D was tested using recombinant vaccinia-based IFN-gamma Elispot assay. 47/74 women had a positive Elispot response to at least one clade, and cross-clade responses were frequent. Clade A responses were most frequent, as expected, as clade A dominates the Kenyan epidemic (85% of the 47 women responded to clade A Env, 74% to B, 62% to C, and 43% to D). Cross-clade CD8 T-cell responses were common and directed at conserved epitopes.
- There was a greater magnitude of response to A clade peptides in individuals who responded to more than 1 clade; a 2-fold higher response was observed in clade A in 36% (9/25) of these individuals, and the response to A peptides was never lower. ERYLRDQQL responses were detected in 2 women who had Env responses to all 4 clades, and clade A gave the highest responses; a ERYLkDQQL variant was in clade B and

C, and the clade D Env carried ERsLkDQQL. The epitope VSGFLALAW was also recognized by 1 of the women.

 HLA-B\*5802 was the only HLA common to both women who reacted with ERYLRDQQL, so may be the presenting allele.

Author Location gp160 (584–594)
Author Location gp41 (584–594)
Epitope ERYLKDQQLLG
Immunogen HIV-1 infection
Species (MHC) human

**Donor MHC** A1, A1, B8, B14, Cw7, Cw8 **Assay type** CD8 T-cell Elispot - IFNγ

**Keywords** binding affinity, acute/early infection, early-expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

HXB2 Location gp160 (585–592)
Author Location gp41 (584–591 SF2)
Epitope RYLRDQQL
Immunogen HIV-1 infection
Species (MHC) human (A\*2402)
References Ikeda-Moore et al. 1997

Defined using reverse immunogenetics – 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) – 53 of the 59 peptides bound A\*2402.

• This peptide induced CTL in 2/4 HIV-1 + people tested.

 RYLRDQQL bound to A\*2402 weakly, the epitope can be processed in a vaccinia construct and presented – two specific CTL clones were obtained.

HXB2 Location gp160 (585–592)

Author Location gp41 (590–597 LAI)

Epitope RYLKDQQL

Subtype B

Immunogen HIV-1 infection

Immunogen HIV-1 infection Species (MHC) human (B27) References Shankar *et al.* 1996

**HXB2 Location** gp160 (585–593)

Author Location gp41 (585–593)
Epitope RYLKDQQLL
Immunogen HIV-1 infection
Species (MHC) human (A\*2301)

Donor MHC A\*2301, B\*3501, B\*1503, Cw2, Cw7

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** acute/early infection, early-expressed proteins **References** Cao *et al.* 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

**HXB2 Location** gp160 (585–593) **Author Location** Env

**Epitope** RYLKDQQLL **Subtype** B

Immunogen HIV-1 infection Species (MHC) human (A\*2301)

**Donor MHC** A\*2301, A\*6801, B\*5801, B\*5802

Country United States.

Keywords escape, acute/early infection

References Bernardin et al. 2005

- Full HIV genomes from nine individuals were analyzed for mutations prior to seroconversion and 7 to 28 days later. It was found that the influence of the host's HLA type was reflected within weeks of infection; a statistically significant number of early nonsynonymous mutations were observed within previously reported CTL epitopes.
- K to E mutation was observed in position 4.

**HXB2 Location** gp160 (585–593)

Author Location gp41 (584-591 SF2)

Epitope RYLRDQQLL

**Immunogen** HIV-1 infection **Species (MHC)** human (A\*2402)

**References** Ikeda-Moore *et al.* 1997

- Defined using reverse immunogenetics 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) 53 of the 59 peptides bound A\*2402.
- $\bullet\,$  This peptide induced CTL in 4/4 HIV-1 + people tested.

 RYLRDQQLL bound to A\*2402 strongly, the epitope can be processed in a vaccinia construct and presented – two specific CTL clones were obtained.

**HXB2 Location** gp160 (585–593) **Author Location** gp41 (591–598 LAI)

Epitope RYLKDQQLL

Subtype B

Immunogen

Species (MHC) human (A\*2402)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*2402 epitope.

**HXB2 Location** gp160 (585–593)

**Author Location** (C consensus)

**Epitope** RYLKDQQLL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** gp160 (585–593)

 $\textbf{Author Location} \ gp41 \ (74\text{--}82)$ 

Epitope RYLKDQQLL

Immunogen HIV-1 infection

Species (MHC) human (A23)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** gp160 (585–593)

**Author Location** gp41

Epitope RYLKDQQLL

**Epitope name** A24-RL9(gp41)

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A24)

**Donor MHC** A24, B7, B27

Keywords HAART, ART, supervised treatment interrup-

tions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals. Patient C displayed the greatest response to B27-KK10(p24), and in decreasing order also responded to A24-RW8(Nef), B7-IL9(gp41), A24-RL9(gp41), A24-YL8(gp41), and B7-TM9(Nef).

**HXB2 Location** gp160 (585–593)

**Author Location** Env

Epitope RYLKDQQLL

Epitope name RW8

Immunogen HIV-1 infection

Species (MHC) human (A24)

**Donor MHC** A2, A24, B38, B60, Cw2, Cw12

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords acute/early infection, early treatment

References Montefiori et al. 2003

• HIV-1 + patient AC10 underwent early HAART treatment, which was discontinued 1.5 years later. At this timepoint potent NAb responses against autologous virus were detected. Treatment interruption initially induced weak CD8+ responses directed against 5 epitopes. By days 873d and 923d the CTL response had broadened to target 22 epitopes; of these six were fully characterized. Eventually the virus escaped the NAb response, but escape was not accompanied by a rise in viral load, and the authors suggest the virus was contained by the CTL response.

**HXB2 Location** gp160 (585–595)

Author Location gp41 (584–591 SF2)

Epitope RYLRDQQLLGI

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

**References** Ikeda-Moore *et al.* 1997

- Defined using reverse immunogenetics 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) 53 of the 59 peptides bound A\*2402.
- This peptide induced CTL in 4/4 HIV-1 + people tested.

• RYLRDQQLLGI bound to A\*2402 with medium strength, the epitope can be processed in a vaccinia construct and presented – two specific CTL clones were obtained.

HXB2 Location gp160 (585–595)
Author Location Env (584–594)
Epitope RYLRDQQLLGI
Epitope name Env584-11
Immunogen vaccine

Vector/Type: Sendai virus vector system

(SeV)

Species (MHC) human (A\*2402)

References Kawana-Tachikawa et al. 2002

- A Sendai virus vector system (SeV) was developd that expressed HLA-A\*2402-restricted class I/peptide complexes; this system could be used to detect responses and has the potential to elicit immune responses.
- MHC class I/peptide tetramers could be made using this system that bound to epitope-specific CTLs in PBMCs.
- Cells transfection with SeV modified to express A\*2402-HIV epitope complexes induced CTL mediated specific cell lysis.

HXB2 Location gp160 (586–593) **Author Location** gp41 (584–591 NL43)

Epitope YLKDQQLL Immunogen HIV-1 infection Species (MHC) human (A\*2402) References Dai et al. 1992

- The lysine (K) is critical for eliciting a HLA-A24 CTL response.
- C. Brander notes that this is an A\*2402 epitope in the 1999 database, and suggested that the epitope is RYLKQQLL.

**HXB2 Location** gp160 (586–593)

**Author Location** gp41 (591–598)

Epitope YLRDQQLL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A24)

**Keywords** subtype comparisons, HIV exposed persistently seronegative (HEPS), immunodomi-

nance

References Kaul et al. 2001a

- Variants (R)YL(R/K)DQQLL are specific for the A/B clade.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-A24 women, 3/4 HEPS and 10/10 HIV-1 infected women recognized this epitope, and (R)YL(R/K)DQQLL tended to be reactive in HEPS and infected women, RDYV-DRFFKTL in infected women only.

- The dominant response to this HLA allele was to this epitope in all 3/4 HEPS cases but in only 4 of the 10/10 HIV-1 infected women.
- Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.

HXB2 Location gp160 (586-593)

Author Location gp41 (580–587 CM243 subtype CRF01)

Epitope YLKDQQLL
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A24)

**Keywords** subtype comparisons

References Bond et al. 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- The only HLA-A24 FSW tested did not recognized the E clade version of this epitope RYLKDQKLL, which differs from the previously defined B clade version by one amino acid, YLKDQQLL, with an additional amino acid added on.

HXB2 Location gp160 (586-593)

Author Location gp41 (591–598)

**Epitope** YLKDQQLL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A24)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- One of seven patients responded to this peptide with GzB producing cells, and a different patient responded with IFNgamma producing cells.

**HXB2 Location** gp160 (586–593)

Author Location gp41 (586-593 LAI)

Epitope YLKDQQLL

Epitope name E1

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A24, B8)

Keywords HAART, ART

References Mollet et al. 2000

 A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.

- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+PBL but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

**HXB2 Location** gp160 (586–593) **Author Location** gp41 (subtype A)

Epitope YLKDQQLL

Subtype A

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (A24, B8)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

**HXB2 Location** gp160 (586–593)

Author Location gp41 (586–593)

Epitope YLKDQQLL

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*0801 epitope.

HXB2 Location gp160 (586-593)

**Author Location** gp41

Epitope YLKDQQLL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

**Donor MHC** A\*0101, A\*0301, B\*0801, B\*5101; A\*0101, B\*0801

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , HLA binding **Keywords** escape, acute/early infection, variant cross-

recognition or cross-neutralization

References Milicic et al. 2005

- CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient. In an early acute infection of the donor, the precise timing of transmission determines the viral variants transmitted.
- The second donor in the study shares A\*0101 and B\*0801 with his partner. Escape variant ylQdqqll was transmitted, and it reduces binding to B\*0801 by 92% relative to YLKDQQLL.

HXB2 Location gp160 (586-593)

**Author Location** gp41 (586–593)

Epitope YLKDQQLL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Johnson et al. 1992

 Two overlapping CTL epitopes were mapped with different HLA restriction (also see ERYLKDQQL HLA-B14)

**HXB2 Location** gp160 (586–593)

**Author Location** gp41 (586–593)

Epitope YLKDQQLL

Immunogen peptide-HLA interaction

Species (MHC) human (B8)

References Sutton et al. 1993

 Predicted epitope based on B8-binding motifs, from larger peptide QLQARILAVERYLKDQQLLGIWGCS.

**HXB2 Location** gp160 (586–593)

**Author Location** gp41 (76–83)

Epitope YLKDQQLL

Immunogen

Species (MHC) human (B8)

References Goulder et al. 1997g

• Included in a study of the B8 binding motif.

**HXB2 Location** gp160 (586–593)

**Author Location** gp41

Epitope YLKDQQLL

**Immunogen** 

Species (MHC) human (B8)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied – these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive.
- HIV-2 sequence: YLQDQARL no cross-reactivity Johnson *et al.* [1992]

HXB2 Location gp160 (586-593)

Author Location gp41 (586–593)

Epitope YLKDQQLL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B8)

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

HXB2 Location gp160 (586-593)

Author Location gp41 (586-593)

Epitope YLKDQQLL
Immunogen HIV-1 infection
Species (MHC) human (B8)
References Day et al. 2001

 B8-restricted CTL accounted for about 1/3 of the total CTL response in one individual.

HXB2 Location gp160 (586-593)

**Author Location** Env (586–593)

Epitope YLKDQQLL

Epitope TERDQQ

Epitope name YL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A\*01, A\*11, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** escape, characterizing CD8+ T cells, optimal

epitope

References Koibuchi et al. 2005

- HIV-1-specific CD8 T-cell responses were shown to be persistent in the chronic phase of HIV-1 infection, although the responses to some of the epitopes were declining despite the persistence of the targeted sequences in vivo. Only 4/14 epitopes were potential CTL escape variants, although strong responses to these epitopes persisted for 6 years. This indicates limited viral evolution within targeted CD8 T-cell epitopes during the chronic phase of infection.
- The autologous form of the epitope, YLKDQQLL, matched the B consensus throughout the 5-year period of study, except for 1 rare variant at the first time point, YLrDQQLL, and 1 at year 5, YLKgQQLL.

HXB2 Location gp160 (586-593)

**Author Location** gp160

Epitope YLRDQQLL

Immunogen HIV-1 infection

Species (MHC) human

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls, ML887.

**HXB2 Location** gp160 (586–598)

Author Location gp41 (586-598)

 ${\bf Epitope} \ {\tt YLRDQQLLGIWGC}$ 

Immunogen HIV-1 infection

Species (MHC) human (Cw7)

References Nehete et al. 1998a

- Three long-term non-progressors and one asymptomatic HIV+ individual were studied and found to have HLA class I Crestricted CD8+ Env-specific CTLs – Cw7 specific CTL were found against three peptides, including this one.
- HLA-C antigens are expressed on lymphoid cells to a lesser extent, 10% of either HLA-A or HLA-B.
- HLA-C confers protection against lysis by natural killer cells and by non-MHC-restricted effector T cells and Cw7 directly governs this resistance to lysis – the authors hypothesize that pathogens that inhibit antigen expression and class I expression may particularly down regulate Cw7, thus triggering non-MHC restricted killing.

HXB2 Location gp160 (594-608)

**Author Location** gp41

Epitope GIWGCSGKLICTTAV

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Jin et al. 1998b

- Progressive HIV-1 infection and CD4 decline was associated decreased the IL-2-expandable HIV-1 CTL response in 10 asymptomatic HIV-infected patients – this observation may be partially due to a reduction and impaired function of T helper cells, CTL exhaustion and APC dysfunction.
- Continued presence of HIV-1 specific memory cells (CTLp) was observed in three patients, one to GIWGCSGKLICTTAV, one to ALIWEDLRSLCLFSY, and one to PIPIHYCAPAG-FAILKCNNK.

**HXB2 Location** gp160 (594–608)

**Author Location** gp41 (SF2)

Epitope GIWGCSGKLICTTAV

Epitope name Peptide2

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

Assay type Chromium-release assay

References Carmichael et al. 1996

Cross-reactivity of Env-specific CTL clones from 14 seropositive HIV-1 infected patients was tested using peptides based on 3 B clade variants (MN, IIIB, and RF). The proportion of CTL clones that cross-recognized conserved variants was low. Most

CTL clones recognized only one peptide variant, indicating most Env responses are not cross-reactive within the B clade.

• This HLA B17(57) epitope was newly identified within gp41 of HIV-1 SF2. SF2 and IIIB have identical sequences within this peptide, but the T-cell clone that recognizes this peptide does not recognize the MN (gFwgcsgklicttTv) or RF (giwgcsgklicttTv) variants of this peptide.

HXB2 Location gp160 (606–614) Author Location gp41 (605–615 LAI)

Epitope TAVPWNASW

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (B\*3501)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*3501 epitope.

**HXB2 Location** gp160 (606–614)

Author Location gp41 (606–614 HXB2)

**Epitope** TAVPWNASW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

Keywords epitope processing

References Ferris et al. 1996

 Natural form of this peptide is not glycosylated, suggesting initial Class I processing may occur in the cytosol.

**HXB2 Location** gp160 (606–614)

Author Location gp41 (605-615 LAI)

Epitope TAVPWNASW

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (B35)

References Johnson et al. 1994b

• Epitope for vaccine induced CD8+ clone.

**HXB2 Location** gp160 (606–614)

Author Location gp41 (606-614 LAI)

**Epitope** TAVPWNASW

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component:

gp160

Species (MHC) human (B35)

References Johnson et al. 1994a

 HLA restricted CTL response to epitope in HIV-1 vaccinia-env vaccinees.

**HXB2 Location** gp160 (606–614)

Author Location gp41 (606-614 LAI)

**Epitope** TAVPWNASW

Subtype B

Immunogen vaccine

Vector/Type: vaccinia HIV component: gp160

Species (MHC) human (B35)

References Hammond et al. 1995

• Peptide only processed by a TAP-1/2-dependent pathway.

HXB2 Location gp160 (606-614)

Author Location gp41 (606-614)

**Epitope** TAVPWNASW

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Ferris et al. 1999

• This epitope is processed by a TAP1/2 dependent mechanism.

**HXB2 Location** gp160 (606–614)

**Author Location** gp41 (subtype B)

**Epitope** TAVPWNASW

Subtype B

**Immunogen** HIV-1 exposed seronegative

Species (MHC) human (B35)

Keywords subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among A, B and D clade viruses.

HXB2 Location gp160 (606-614)

Author Location gp41 (606–614)

**Epitope** TAVPWNASW

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B35)

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

HXB2 Location gp160 (606–614)

Author Location gp41 (606–614)

**Epitope** TAVPWNASW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

**Donor MHC** A3, A33, B14, B35, Cw\*0401, Cw\*0802

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

**HXB2 Location** gp160 (606–614)

**Author Location** Env (96–104)

**Epitope** TAVPWNASW

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/9 patients recognized this epitope.

**HXB2 Location** gp160 (614–631)

Author Location (C consensus)

Epitope WSNKSQEEIWDNMTWMQW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

HXB2 Location gp160 (614-631)

Author Location (C consensus)

Epitope WSNKSQEEIWDNMTWMQW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0401)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Kiepiela et al. 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
 Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** gp160 (634–648)

**Author Location** gp41 (641–655 SF2)

Epitope EIDNYTNTIYTLLEE

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-A1, A2, B51, and B57.

**HXB2 Location** gp160 (678–686)

**Author Location** Env (679–687 subtype B)

Epitope WLWYIKIFI

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human (A\*0201)

**Keywords** binding affinity

References Kundu et al. 1998a

- Ten HIV-1 + HLA A2 asymptomatic individuals were given two courses of HIV-1 MN rgp160 vaccine over a 2 year period.
- Two hundred and fifty three HIV-1 peptides of 9 or 10 aa possessing the HLA-A2.1 binding motif (Leu at position 2, Val at the C terminus) were identified in gp160, of which 25 had a high or intermediate binding affinity.
- Eleven peptides were studied that had high HLA-A2 binding affinity a CTL response was detected to 9/11 peptides in at least 1 individual.
- CTL responses after reimmunization may include recall responses only individuals with vaccine cross-reactive sequences prior to vaccination showed detectable CTL responses.

**HXB2 Location** gp160 (680–688)

Author Location gp41 (679-687 SF2)

Epitope WYIKIFIMI

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

References Ikeda-Moore et al. 1997

Defined using reverse immunogenetics – 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) – 53 of the 59 peptides bound A\*2402.

• This peptide induced CTL in 1/4 HIV-1 + people tested.

 WYIKIFIFMI bound to A\*2402 strongly, the epitope can be processed in a vaccinia construct and presented – two specific CTL clones were obtained.

HXB2 Location gp160 (681-689)

**Author Location** Env (681–)

**Epitope** YIKIFIMIV

Epitope name Env681

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Env Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (MHC) human, transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a good A2 binder, and induced CTL and CD8+ T-cell IFN gamma responses in mice. Responses were detected in 2/17 HIV+ HLA-A2 subjects.

HXB2 Location gp160 (685–693)

Author Location Env (686–694 subtype B)

Epitope FIMIVGGLV

Subtype B

Immunogen vaccine

*Vector/Type:* protein *Strain:* B clade MN *HIV component:* gp160

Species (MHC) human (A\*0201)

**Keywords** binding affinity

References Kundu et al. 1998a

- Ten HIV-1 + HLA A2 asymptomatic individuals were given two courses of HIV-1 MN rgp160 vaccine over a 2 year period.
- Two hundred and fifty three HIV-1 peptides of 9 or 10 aa possessing the HLA-A2.1 binding motif (Leu at position 2, Val at the C terminus) were identified in gp160, of which 25 had a high or intermediate binding affinity.
- Eleven peptides were studied that had high HLA-A2 binding affinity a CTL response was detected to 9/11 peptides in at least 1 individual.
- CTL responses after reimmunization may include recall responses only individuals with vaccine cross-reactive sequences prior to vaccination showed detectable CTL responses.
- ALTERNATIVE EPITOPE: IMIVGGLVGL no CTL response was shown to the peptides FIMIVGGLV or IMIVGGLVGL.

HXB2 Location gp160 (698–707)

**Author Location** Env (696–706)

Epitope VFAVLSIVNR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*3303)

References Hossain et al. 2001; Takiguchi et al. 2000

- HLA-A33 a very common allele in Asian, with HLA-A\*3303 the most common among the Japanese. New A\*3303 epitopes were defined to better characterize the immune response in this population.
- The anchor motif for HLA\*3303 (A, I, L, V, F, Y in position 2 (F and Y bind most strongly), and R (K is also tolerated) in the C-terminal position) was used to define 82 potentially reactive peptides in Env; 37/82 peptides bound to A\*3303; 3/37 peptides could induce peptide-specific CTL in bulk PBMC cultures from 1/3 HLA A\*3303 positive individuals tested.
- CTL clones were isolated that killed target cells in a concentration dependent manner after pulsing with the VFAVLSIVNR peptide, that could also kill cells transfected with env expressed from a vaccinia vector. Bulk cultures were tested from six additional people, and only 1/6 reacted with this peptide, but the peptide is in a highly variable region.

HXB2 Location gp160 (698–707)

**Author Location** gp41 (187–196)

**Epitope** VFAVLSIVNR

Immunogen HIV-1 infection

Species (MHC) human (A\*3303)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location gp160 (700–708)

Author Location gp41 (705–714)

Epitope AVLSVVNRV

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Ferris et al. 1999

• This epitope is processed by a TAP1/2 dependent mechanism.

**HXB2 Location** gp160 (700–708)

Author Location Env (695-705 BH10, LAI)

Epitope AVLSVVNRV

Immunogen HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is LRIVFAVLSVV) has similarity with the human chemokine-factor 3 fragment LRLVFALVTAV.

HXB2 Location gp160 (701–719)

**Author Location** Env (691–710)

Epitope VLSIVNQVRRQGYSPLSFQT

Immunogen HIV-1 infection

Species (MHC) human (B15)

**Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4, DR8, DO7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 5 epitopes from this individual that varied over time where the internal mutations became fixed; a dramatic decrease in CTL activity against the wild type epitope was observed as the mutation arose. The vlsivnKvrrqgysplsfqt variant found at 20 and 47 months postseroconversion.

HXB2 Location gp160 (701–720) Author Location gp41 (701–720 BH10) Epitope VLSIVNRVRQGYSPLSFQTH

Immunogen HIV-1 infection Species (MHC) human (A32) References Safrit *et al.* 1994a

· Recognized by CTL derived from acute seroconverter.

**HXB2 Location** gp160 (702–721) **Author Location** Env (702–721)

Epitope LSIVNRVRQGYSPLSFQTLT

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** gp160 (703–712) **Author Location** gp160 (703–712)

Epitope EIIFDIRQAY

Epitope name EY10

Immunogen HIV-1 infection

**Species (MHC)** human (A\*2501) **Keywords** optimal epitope

**References** Frahm *et al.* 2007

• C. Brander notes this is a A\*2501 epitope.

HXB2 Location gp160 (704–712)

**Author Location** gp160 (704–712 LAI)

Epitope IVNRNRQGY

Subtype B

Immunogen

Species (MHC) human (A\*3002)

**Keywords** optimal epitope

References Frahm et al. 2007; Goulder et al. 2001a

• C. Brander notes this is an A\*3002 epitope.

HXB2 Location gp160 (704–712)

Author Location gp41

Epitope IVNRVRQGY

Epitope name IY9 (gp41)

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

References Goulder et al. 2001a

- HLA-A\*3002 is very common in African populations, 50% of Zimbabweans express HLA-A30, 44% in African Zulu, so five new HIV epitopes were characterized that are presented by this HLA molecule.
- A rapid method was developed combining ELISPOT with intracellular IFN-γ staining of PBMCs to map optimal epitopes, then HLA presenting molecules were defined this method was completed within 48 to 72 hours of receipt of blood.
- Two individuals were studied: Subject 199 (HLA A\*0201/\*3002 B\*4402/51 Cw2/5), a Caucasian, and Subject 6007 (HLA A\*3002/ B53/\*5801 Cw4/7) an African-Caribbean.
- In both HLA-A\*3002 individuals the response to RSLYNT-VATLY was dominant.
- In subject 199 four additional A\*3002 epitopes were identified.
- Three quantitative assays, ELISPOT, precursor frequency and chromium release, confirmed a hierarchy of response: RY11 (p17) > KY9 (gp41) > KY9 (RT-53) > IY9 (gp41)

HXB2 Location gp160 (712–720)

Author Location gp41 (201–209)

Epitope YSPLSLQTL

Epitope name YL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0102)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101, Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** escape, immune evasion, optimal epitope, HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- This is a newly defined epitope. Last position (9)in the epitope had potentially experienced positive selection. YSPLSLQTr escape variant was found.

**HXB2 Location** gp160 (742–761)

**Author Location** Env (742–761)

Epitope RDRSIRLVSGFLALAWDDLR

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.

This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.
 A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.

HXB2 Location gp160 (747–755) Author Location gp41 (747–755) Epitope RLVNGSLAL Immunogen HIV-1 infection Species (MHC) human (A2) References Parker *et al.* 1992

• Studied in the context of HLA-A2 peptide binding.

**HXB2 Location** gp160 (747–755)

**Author Location** gp41 (741–749 CM243 subtype CRF01)

Epitope RLVSGFLAL
Epitope name E747-755
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A2)

Keywords HIV exposed persistently seronegative

(HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was reactive in HIV+ control study subject 144 who carried HLA-A2.

**HXB2 Location** gp160 (747–755)

**Author Location** gp41 (741–749 CM243 subtype CRF01)

Epitope RLVSGFLAL
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A2)
Keywords subtype comparisons

**References** Bond *et al.* 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- 2/4 tested FSWs recognized the E clade version of this epitope, which differs from the previously defined B clade version by two amino acids, RLVNGSLAL.
- This epitope was somewhat conserved 4/8 subtypes: CRF01 (E), B, C, and G.

**HXB2 Location** gp160 (747–763)

Author Location (C consensus)

Epitope RLVSGFLALAWDDLRSL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0202)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** gp160 (749–757)

**Author Location** gp160

Epitope VSGFLALAW Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: vaccinia Strain: A clade, B clade, D clade NDK, C clade consensus HIV

component: Env

Species (MHC) human

**Donor MHC** A\*0202, A\*7401, B\*5802, B\*1503,

Cw\*0202, Cw\*0602

Country Kenya.

Assay type CD8 T-cell Elispot - IFNγ, Other

**Keywords** subtype comparisons, variant cross-

recognition or cross-neutralization

References McKinnon et al. 2005

- Interclade cross-reactivity of clades A, B, C, and D was tested using recombinant vaccinia-based IFN-gamma Elispot assay. 47/74 women had a positive Elispot response to at least one clade, and cross-clade responses were frequent. Clade A responses were most frequent, as expected, as clade A dominates the Kenyan epidemic (85% of the 47 women responded to clade A Env, 74% to B, 62% to C, and 43% to D). Cross-clade CD8 T-cell responses were common and directed at conserved epitopes.
- There was a greater magnitude of response to A clade peptides in individuals who responded to more than 1 clade; a 2-fold higher response was observed in clade A in 36% (9/25) of these individuals, and the response to A peptides was never lower. ERYLRDQQL responses were detected in 2 women who had Env responses to all 4 clades, and clade A gave the highest responses; an ERYLkDQQL variant was in clade B and C, and the clade D Env carried ERsLkDQQL. The epitope VSGFLALAW was also recognized by 1 of the women; A and C clade were identical, while clade B carried VnGsLiLAW, clade D VnGlsAiAW.

**HXB2 Location** gp160 (754–768)

**Author Location** gp41

Epitope ALIWEDLRSLCLFSY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B55)

References Jin et al. 1998b

- Progressive HIV-1 infection and CD4 decline was associated decreased the IL-2-expandable HIV-1 CTL response in 10 asymptomatic HIV-infected patients – this observation may be partially due to a reduction and impaired function of T helper cells, CTL exhaustion and APC dysfunction.
- Continued presence of HIV-1 specific memory cells (CTLp) was observed in three patients, one to GIWGCSGKLICTTAV, one to ALIWEDLRSLCLFSY, and one to PIPIHYCAPAG-FAILKCNNK.

**HXB2 Location** gp160 (754–768)

Author Location gp41 (SF2)

**Epitope** ALIWERDLRSLCLFSY

Epitope name Peptide78

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B55)

**Assay type** Chromium-release assay **References** Carmichael *et al.* 1996

- Cross-reactivity of Env-specific CTL clones from 14 seropositive HIV-1 infected patients was tested using peptides based on 3 B clade variants (MN, IIIB, and RF). The proportion of CTL clones that cross-recognized conserved variants was low. Most CTL clones recognized only one peptide variant, indicating most Env responses are not cross-reactive within the B clade.
- This HLA B22(55) epitope was defined using SF2 peptides.
   The CTL clone that recognized it did not cross-recognize the MN, IIIB, or RF variants of this peptide.

**HXB2 Location** gp160 (760–767)

Author Location gp41 (760–767)

**Epitope** LRSLFLFS

Immunogen HIV-1 infection

Species (MHC) human (A\*2301)

Donor MHC A\*2301, B\*3501, B\*1503, Cw2, Cw7

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** binding affinity, acute/early infection, early-expressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

**HXB2 Location** gp160 (767–775)

Author Location gp41 (766–774 SF2)

Epitope SYRRLRDLL

Immunogen HIV-1 infection Species (MHC) human (A\*2402)

**References** Ikeda-Moore *et al.* 1997

• Defined using reverse immunogenetics – 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) – 53 of the 59 peptides bound A\*2402.

- This peptide induced CTL in 1/4 HIV-1 + people tested.
- SYRRLRDLL bound to A\*2402 moderately, the epitope can be processed in a vaccinia construct and presented – two specific CTL clones were obtained.

**HXB2 Location** gp160 (767–780)

Author Location gp41 (606-614 LAI)

**Epitope** SYHRLRDLLLIVTR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A31)

References Hammond et al. 1995

- Peptide only processed by a TAP-1/2-dependent pathway.
- CTL from an acute seroconverter.

**HXB2 Location** gp160 (769–777)

Author Location gp41 (769–777 BH10)

**Epitope** HRLRDLLLI

Immunogen HIV-1 infection

Species (MHC) human

References Safrit et al. 1994a

· Recognized by CTL derived from acute seroconverter.

**HXB2 Location** gp160 (770–778)

Author Location Env (679–777)

Epitope RLRDLLLIV

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords binding affinity

References Kmieciak et al. 1998a

- CTL responses in six patients to four Env epitopes were studied: D2: LLNATAIAV, 5.3: RLRDLLLIV, D1: KLTPLCVTL, and 4.3: QMHEDIISL all have A2 anchor residues.
- The C terminal epitopes (D2 and 5.3) were highly variable and the variability was considered responsible for limited CTL response, while D1 and 4.3, N-terminal epitopes, were much more conserved and gave evidence of high levels of CTL response *in vitro*.
- Peptides 5.3 and D2 bound to HLA A\*0201 with low affinity.

**HXB2 Location** gp160 (770–780)

Author Location gp41 (768–778 NL43)

Epitope RLRDLLLIVTR

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

References Takahashi et al. 1991

• CD8+ T cell clone.

**HXB2 Location** gp160 (770–780)

Author Location gp41 (775–785 LAI)

Epitope RLRDLLLIVTR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0301 epitope.

HXB2 Location gp160 (770-780) **Author Location** gp41 (770–780 BH10) Epitope RLRDLLLIVTR Immunogen HIV-1 infection

Species (MHC) human (A\*3101)

References Safrit et al. 1994a; Safrit et al. 1994b

- · Recognized by CTL derived from acute seroconverter.
- C. Brander notes that this is an A\*3101 epitope in the 1999 database.

HXB2 Location gp160 (770–780) Author Location gp160 (770–780 LAI)

Epitope RLRDLLLIVTR

Subtype B

**Immunogen** 

Species (MHC) human (A\*3101)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is an A\*3002 epitope.

HXB2 Location gp160 (770–780) Author Location gp41 (768–778 NL43) Epitope RLRDLLLIVTR Immunogen HIV-1 infection

Species (MHC) human (A3)

**Keywords** subtype comparisons References Cao et al. 1997a

- The consensus peptide of clade B is RLRDLLLIVTR.
- The consensus peptide of clades A, C and E is RLRDFILIVTR and it is less reactive.
- The consensus peptide of clade D is SLRDLLLIVTR and it is less reactive.

HXB2 Location gp160 (770-780) Author Location gp41 (775–785)

Epitope RLRDLLLIVTR

Immunogen HIV-1 infection, HIV-1 exposed seronegative

**Species (MHC)** human (A3)

Keywords HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

• ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

HXB2 Location gp160 (770-780)

Author Location gp41 (770–780)

Epitope RLRDLLLIVTR Immunogen HIV-1 infection Species (MHC) human (A3)

Keywords rate of progression, acute/early infection

References Day et al. 2001

• The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)

- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- All patients recognized at least 1 A3 epitope, up to 8 A3 epitopes, but none was clearly dominant.

HXB2 Location gp160 (770-780)

**Author Location** Nef (73–82)

Epitope RLRDLLLIVTR

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- All patients recognized at least 1 A3 epitope, up to 8 A3 epitopes, but none was clearly dominant.
- In two of the subjects, RLRDLLLIVTR was the dominant epitope.

**HXB2 Location** gp160 (770–780)

Author Location gp41 (769–780)

Epitope RLRDLLLIVTR

Epitope name A3-RR11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 1/7 individuals began to have detectable responses to this epitope after STI.

HXB2 Location gp160 (770–780)

**Author Location** Env (770–780)

Epitope RLRDLLLIVTR

Immunogen HIV-1 infection

Species (MHC) human (A3)

DR8, DO7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 7 epitopes from this individual that varied over time, although the internal mutations did not become fixed. rlrdlllVItr variant residues were found. The V mutation arose at late time points, the I mutation arose at intermediate time points.

HXB2 Location gp160 (770-780)

**Author Location** Env (786–778)

Epitope RLRDLLLIVTR

Immunogen HIV-1 infection

Species (MHC) human (A3)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 14 patients recognized this epitope.

HXB2 Location gp160 (770-780)

Author Location gp41 (770–780)

Epitope RLRDLLLIVTR

Immunogen HIV-1 infection

Species (MHC) human (A31)

References Ferris et al. 1999; Hammond et al. 1995

• This epitope is processed by a TAP1/2 dependent mechanism.

**HXB2 Location** gp160 (770–780)

**Author Location** gp41 (770–780)

Epitope RLRDLLLIVTR

Immunogen HIV-1 infection

Species (MHC) human (A31)

**Donor MHC** A\*0201, A31, B44, B60, Cw3, Cw16

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- **Donor MHC** A03, A32, B51, B15, Cw03, Cw06, DR4, CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFNsecreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
  - · All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
  - More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

HXB2 Location gp160 (770–780)

**Author Location** Env

Epitope RLRDLLLIVTR

Epitope name TW10

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A31)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assav

Keywords epitope processing, escape

References Draenert et al. 2004b

• This study characterizes the N-terminal flanking position of the epitope ISPRTLNAW, and mutations in this position are thought to impact processing. The A31 epitope RLRDLL-LIVTR was used as a negative control in this study.

HXB2 Location gp160 (770–780)

Author Location gp41 (775–785)

Epitope RLRDLLLIVTR

Immunogen HIV-1 infection

Species (MHC) human

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the HLA A2+ was HLA A\*0201, A31, B8, B51 and responded to this epitope that has been previously noted to be HLA A3.1, as well as seven others.

HXB2 Location gp160 (777–785)

Author Location gp41 (782–790 LAI)

**Epitope IVTRIVELL** 

Subtype B

Immunogen

Species (MHC) human (A\*6802)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*6802 epitope.

HXB2 Location gp160 (781-802) Author Location gp41 (788–809 HXB2)

Epitope IVELLGRRGWEALKYWWNLLQY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B27)

References Lieberman et al. 1992

• CTL epitope defined by T cell line and peptide mapping.

HXB2 Location gp160 (781–802) Author Location gp120 (788-809)

Epitope IVELLGRRGWEALKYWWNLLQY

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

• HIV-specific CTL lines developed by ex vivo stimulation with peptide.

HXB2 Location gp160 (786-794)

Author Location gp41 (751–759)

**Epitope** GRRGWEALK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*2705)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, CD8 T-cell Elispot

granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- One of seven patients responded to this peptide with GzB producing cells, and a different patient responded with IFNgamma producing cells.

HXB2 Location gp160 (786-794)

Author Location gp41 (791-799 LAI)

Epitope GRRGWEALK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords review

References McMichael & Walker 1994

- Review of HIV CTL epitopes.
- Also: J. Liebermann 1992 and pers. comm. J. Liebermann.

**HXB2 Location** gp160 (786–795)

Author Location gp41 (791–800 LAI)

Epitope GRRGWEALKY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*2705)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*2705 epitope.

**HXB2 Location** gp160 (786–795)

Author Location gp41 (791–800 LAI)

Epitope GRRGWEALKY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Lieberman 1998

• Optimal peptide mapped by titration J. Lieberman, pers. comm.

HXB2 Location gp160 (786–795)

Author Location gp41 (786–795)

**Epitope** GRRGWEALKY

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Day et al. 2001

**HXB2 Location** gp160 (787–795)

**Author Location** gp160 (787–795)

Epitope RRGWEVLKY

Immunogen HIV-1 infection

Species (MHC) human (A\*0101)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location gp160 (787–795)

**Author Location** gp41 (787–795)

**Epitope** RRGWEVLKY

Immunogen HIV-1 infection

Species (MHC) human (A1)

**Donor MHC** A1, A1, B8, B14, Cw7, Cw8

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFNsecreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

HXB2 Location gp160 (787–805)

**Author Location** (C consensus)

Epitope QRGWEALKYLGSLVQYWGL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

 A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses

**HXB2 Location** gp160 (794–802)

Author Location gp160 (794–802 LAI)

were associated with higher viremia.

**Epitope** KYCWNLLQY

Subtype B

Immunogen

Species (MHC) human (A\*3002)

Keywords optimal epitope

References Frahm et al. 2007; Goulder et al. 2001a

• C. Brander notes this is an A\*3002 epitope.

HXB2 Location gp160 (794–802)

**Author Location** gp41

**Epitope** KYCWNLLQY

Epitope name KY9 (gp41)

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

References Goulder et al. 2001a

- HLA-A\*3002 is very common in African populations, 50% of Zimbabweans express HLA-A30, 44% in African Zulu, so five new HIV epitopes were characterized that are presented by this HLA molecule.
- A rapid method was developed combining ELISPOT with intracellular IFN-γ staining of PBMCs to map optimal epitopes, then HLA presenting molecules were defined this method was completed within 48 to 72 hours of receipt of blood.
- Two individuals were studied: Subject 199 (HLA A\*0201/\*3002 B\*4402/51 Cw2/5), a Caucasian, and Subject 6007 (HLA A\*3002/ B53/\*5801 Cw4/7) an African-Caribbean.
- In both HLA-A\*3002 individuals the response to RSLYNT-VATLY was dominant.
- In subject 199 four additional A\*3002 epitopes were identified.
- Three quantitative assays, ELISPOT, precursor frequency and chromium release, confirmed a hierarchy of response: RY11 (p17) > KY9 (gp41) > KY9 (RT-53) > IY9 (gp41)

HXB2 Location gp160 (794-802)

Author Location gp41 (283-291)

**Epitope** KYCWNLLQY

Immunogen HIV-1 infection

Species (MHC) human (A\*3002)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location gp160 (794-814)

**Author Location** gp41 (SF2)

Epitope KYCWNLLQYWSQELKNSAVSL

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.
- The response to the peptide was CD8 dependent, but the HLA
  presenting molecule and optimal epitope were not determined.

HXB2 Location gp160 (795–816)

Author Location gp41 (802–823 HXB2)

Epitope YWWNLLQYWSQELKNSAVNLLN

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1992

• CTL epitope defined by T cell line and peptide mapping.

HXB2 Location gp160 (799–807)

**Author Location** Env (800–808 subtype B)

Epitope LLQYWSQEL

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human (A\*0201)

Keywords binding affinity

References Kundu et al. 1998a

- Ten HIV-1 + HLA A2 asymptomatic individuals were given two courses of HIV-1 MN rgp160 vaccine over a 2 year period.
- Two hundred and fifty three HIV-1 peptides of 9 or 10 aa possessing the HLA-A2.1 binding motif (Leu at position 2, Val at the C terminus) were identified in gp160, of which 25 had a high or intermediate binding affinity.
- Eleven peptides were studied that had high HLA-A2 binding affinity a CTL response was detected to 9/11 peptides in at least 1 individual.
- CTL responses after reimmunization may include recall responses only individuals with vaccine cross-reactive sequences prior to vaccination showed detectable CTL responses.

**HXB2 Location** gp160 (805–814)

Author Location gp41 (810–819 LAI)

Epitope QELKNSAVSL

Subtype B

Immunogen

Species (MHC) human (B\*4001)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*4001,B60 epitope.

HXB2 Location gp160 (805–814)

**Author Location** gp41 (SF2)

Epitope QELKNSAVSL

**Immunogen** HIV-1 infection

Species (MHC) human (B60)

References Altfeld et al. 2000

 This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes. • B60 is present in 10-20% of the Caucasoid and very common in Asian populations.

• Two hundred and fifty three HIV-1 peptides of 9 or 10 aa possessing the HLA-A2.1 binding motif (Leu at position 2, Val

HXB2 Location gp160 (805–814)
Author Location gp41 (805–814)
Epitope QELKNSAVSL
Immunogen HIV-1 infection
Species (MHC) human (B60, B61)
Keywords immunodominance
References Day et al. 2001

- No immunodominant responses were detected to five B61-restricted epitopes tested.
- All five B60-restricted epitopes were reactive in another subject, and the B60-restricted responses together contributed over onethird of the total CTL response.

**HXB2 Location** gp160 (805–814)

Author Location Env (799-813 BH10, LAI)

**Epitope** QELKNSAVSL **Immunogen** HIV-1 infection

Species (MHC) human

References Maksiutov et al. 2002

- This study employs an antigenic similarity matrix to compare HIV-1 antigenic determinants with human proteins.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is LLQY-WSQELKNSAVS) has similarity with the complement component C6 fragment LTQFSSEELKNSGLT.
- This CTL epitope (the HIV-1 LAI fragment with high similarity to a human protein overlapping this epitope is NSAVSLL-NATAIAVA) also has similarity with the human INT-2 proto-oncogene protein precursor (fibroblast growth factor-3) fragment NSAYSILEITAVEVG.

**HXB2 Location** gp160 (813–822)

Author Location gp41 (814–823 LAI)

**Epitope** SLLNATDIAV

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade MN HIV component: gp160

Species (MHC) human (A\*0201)

References Dupuis et al. 1995

- Of two CTL clones, one reacted only with 815-823, the other with 814-823 and 815-823.
- Noted to be A\*0201 in Brander et al., 1999 database.

**HXB2 Location** gp160 (813–822)

Author Location Env (814–823 subtype B)

**Epitope** SLLNATDIAV

Subtype B

Immunogen vaccine

*Vector/Type:* protein *Strain:* B clade MN *HIV component:* gp160

Species (MHC) human (A\*0201)

Keywords binding affinity

References Kundu et al. 1998a

 Ten HIV-1 + HLA A2 asymptomatic individuals were given two courses of HIV-1 MN rgp160 vaccine over a 2 year period.

- Two hundred and fifty three HIV-1 peptides of 9 or 10 aa possessing the HLA-A2.1 binding motif (Leu at position 2, Val at the C terminus) were identified in gp160, of which 25 had a high or intermediate binding affinity.
- Eleven peptides were studied that had high HLA-A2 binding affinity – a CTL response was detected to 9/11 peptides in at least 1 individual.
- CTL responses after reimmunization may include recall responses only individuals with vaccine cross-reactive sequences prior to vaccination showed detectable CTL responses.
- CTL to overlapping peptides in this region gave a positive response in the greatest number of patients.
- ALTERNATIVE EPITOPES: LLNATDIAV and LLNATDI-AVA – CTL were induced by vaccine in those that had the sequence SLLNATAIAVA in their own infection, but not in those with: NLLNTIAIAVA or NLFNTTAIAVA or SLLNATAITVA.

HXB2 Location gp160 (813–822)

Author Location gp41 (818–827 LAI)

**Epitope** SLLNATDIAV

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human (A\*0201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0201 epitope.

**HXB2 Location** gp160 (813–822)

Author Location gp41 (814–823 LAI)

**Epitope** SLLNATDIAV

Epitope name LR27

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant

(IFA), Montanide (ISA 720), P30, PLG

Species (MHC) mouse (A\*0201)

Keywords binding affinity, vaccine-specific epitope char-

acteristics, immunodominance

References Peter et al. 2001

- The stability of peptide binding to HLA-A2.1 was determined for six HLA-A2.1 peptides included in this vaccine study ILKEPVHGV (RT), SLYNTVATL (p17), SLLNATDIAV (gp41) and LLWKGEGAV (RT) all bound with high affinity comparable to a influenza epitope reference (GILGFVFTL), while RGPGRAFVTI and VIYQYMDDL bound with a lower affinity (relative binding activity = 0.01).
- The four high-affinity peptides formed stable complexes with half-lives ranging between 8 and 32 hours, while the low affinity peptides had half lives of less than an hour.
- HLA-A2.1 transgenic mice were immunized with the six HIV-1 peptides and P30, as a universal T-helper epitope, with IFA or Montanide or microspheres as adjuvants.
- All peptides except VIYQYMDDL induced a stong CTL response in Cr-release assays stronger responses were observed when peptides were delivered alone, indicating immunodominance when the combination was used.

**HXB2 Location** gp160 (813–822) **Author Location** gp41 (814–823 LAI)

**Epitope** SLLNATDIAV **Epitope** name LR27

Subtype B
Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI Adjuvant: Incomplete Freund's Adjuvant (IFA), IL-12, P30

Species (MHC) mouse (A\*0201)

**Keywords** vaccine-specific epitope characteristics, immunodominance

References Peter et al. 2002

• When HIV-1 peptides were used to vaccinate HLA-A2.1 transgenic A2-Kb mice, strong responses to five peptides were observed when the peptides were given individually, but immunodominance limited the response to some of the peptides when they were given in combination Peter et al. [2001]. IL-12 can counteract immunodominance in BALB/c mice, so it was given with the multiple epitope vaccination, and was instead found to specifically eliminate the HLA-A2.1-epitope CTL responses, but not Kb CTL responses. This was possibly a consequence of transient depletion of T-cells, B cells and macropahges in the spleen.

 $\textbf{HXB2 Location} \hspace{0.1cm} gp160 \hspace{0.1cm} (813\text{--}822)$ 

Author Location gp41 (814–823)

Epitope SLLNATDIAV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords dendritic cells

References Kundu et al. 1998b

- Allogeneic dendritic cells (DCs) were obtained from HLAidentical siblings, pulsed with rgp160 MN or A2-restricted HIV-1 epitope peptides, and infused monthly into six HIVinfected patients.
- 1/6 showed increased env-specific CTL and increased lymphoproliferative responses, 2/6 showed increase only in proliferative responses, and 3/6 showed no change – pulsed DCs were well tolerated.
- SLLNATDIAV is a conserved HLA-A2 epitope included in this study – 4/6 patients had this sequence as their HIV direct sequence, and 3 of these had a detectable CTL response – the other two had either the sequence SLFNAIDIAV or SLL-NTTDIVV and no detectable CTL response.
- CTL demonstrated against peptide-coated target, epitope is naturally processed and enhancible with vaccine.

HXB2 Location gp160 (813-822)

Author Location gp41 (818–827)

**Epitope** SLLNATDIAV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.

• 1/11 of the A2+ individuals that didn't respond to SLYNTVATL reacted with seven other epitopes, including this epitope.

**HXB2 Location** gp160 (813–822)

**Author Location** gp41 (SF2)

**Epitope** SLLNATAIAV

Epitope name SV10

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords acute/early infection

References Goulder et al. 2001a

- Dominant CTL epitope in acute infection of patient AC13– response to this epitope corresponded to reduction of initial viremia.
- Several other subdominant CTL epitopes were identified in the acute phase, but a response to SL9, SLYNTVATL, was not evident until 18 months post-presentation.

**HXB2 Location** gp160 (813–822)

**Author Location** gp41 (77–85 SF2)

**Epitope** SLLNATDIAV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A2+ individuals that had a CTL response to this epitope broken down by group: 1/10 group 1, 2/6 group 2, and 1/4 group 3.

HXB2 Location gp160 (813–822)

Author Location gp41 (814–823 CM243 subtype CRF01)

Epitope SLLNATAIAV

Epitope name E813-82

Subtype CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.

• This epitope was reactive in HIV+ control study subjects 125 • Greater resistance was conferred by the gp160deltaV3 than the and 144 who carried HLA-A2.

**HXB2 Location** gp160 (813–822)

Author Location gp41 (814-823 CM243)

Epitope SLLNATAIAV
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A2)
Keywords subtype comparisons

**Keywords** subtype comparisons **References** Bond *et al.* 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- 1/4 tested FSWs recognized the E clade version of this epitope, which differs from the previously defined B clade version by one amino acid, SLLNATDIAV.
- This epitope was somewhat conserved 4/8 subtypes: CRF01 (E), B, D, and F.

**HXB2 Location** gp160 (813–822) **Author Location** gp41 (813–822)

Epitope SLLNATDIAV

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.

**HXB2 Location** gp160 (813–822)

Author Location gp41 (813–822 IIIB)

Epitope SLLNATAIAV

Epitope name D2

Subtype B

Immunogen vaccine

Vector/Type: DNA, DNA with protein boost Strain: B clade IIIB HIV component:

gp160, gp160ΔV3 Adjuvant: IL-12

Species (MHC) mouse (A2)

Keywords vaccine-specific epitope characteristics

References Kiszka et al. 2002

• Transgenic mice expressing a HLA-A2/Kb chimeric protein were vaccinated with a full length gp160 or with gp160deltaV3, with the V3 loop deleted. Mice given gp160deltaV3 had a broader immune response than those given gp160, with increased responses to conserved HLA-A2 epitopes in the C1 region of gp120, KLTPLCVTL, and the C-term region of gp41, SLLNATAIAV.

 Greater resistance was conferred by the gp160deltaV3 than the gp160 vaccine to a challenge of vaccinia expressing heterologous gp160 from primary isolates (VI-06 and 89.6), and the resistance was conferred by CD8+ T-cells.

**HXB2 Location** gp160 (813–822)

Author Location Env (813-)

Epitope SLLNATDIAV

**Epitope name** Env813

Immunogen HIV-1 infection

Species (MHC) human (A2)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This epitope was one of the previously identified HLA-A2 epitopes studied.
- None of the 17 HIV-infected HLA-A2+ people in this study recognized this epitope.

**HXB2 Location** gp160 (813–822)

**Author Location** gp160 (813–822)

**Epitope** SLLNATDIAV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, HLA binding

**Keywords** acute/early infection, optimal epitope

References Altfeld et al. 2005

 The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was recognized both in acute and chronic infection, but slightly more frequently in chronic infection.

HXB2 Location gp160 (813–822)

**Author Location** gp41

Epitope SLLNATDIAV

Epitope name gp41 SV10

Immunogen HIV-1 infection

Species (MHC) human (A68)

**Keywords** binding affinity, subtype comparisons, supertype, computational epitope prediction

References Altfeld et al. 2001c

• HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences – 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 – 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.

- Three additional previously described HLA-A2 epitopes were added to the set of 20, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- This epitope binds to three HLA-A2 supertype alleles: A\*6802 (highest affinity), A\*0202 and A\*0203 (but not A\*0201 and not A\*0206)
- This epitope did not elicit an ELISPOT response in 22 chronic HIV HLA-A2 infections, but elicited a strong response in 1/12 acute HLA-A2 infections this individual, AC13, was HLA A\*0201/68 B44/14 and also had a strong response to HLA-A2 vpr epitope AIIRILQQL.

HXB2 Location gp160 (813–828)

**Author Location** gp41 (MN)

**Epitope** SLLNATAIAVAEGTDR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A2

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords assay standardization/improvement, HAART,

**ART** 

References Chitnis et al. 2003

• 17 perinatally HIV-1 infected children (0.08-16 years) were evaluated for HLA-A2-restricted IFN-gamma CD8+ CTL responses against 4 immunodominant peptides that carry HLA-A2 epitopes. Two peptides were from gp120 (one at position 112, one from the V3 loop), and one each was from gp41 and Gag. 15/17 patients responded to the Gag peptide, 13/17 to the gp41 and the non-V3 gp120 peptides, and 11/17 responded to the V3 loop. 4 children recognized all 4 peptides.

HXB2 Location gp160 (814-822)

**Author Location** Env (815–823)

**Epitope** LLNATAIAV

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

**Keywords** binding affinity

References Kmieciak et al. 1998a

- CTL responses in six patients to four Env epitopes were studied: D2: LLNATAIAV, 5.3: RLRDLLLIV, D1: KLTPLCVTL, and 4.3: QMHEDIISL—all have A2 anchor residues.
- The C terminal epitopes (D2 and 5.3) were highly variable and the variability was considered responsible for limited CTL response, while D1 and 4.3, N-terminal epitopes, were much more conserved and gave evidence of high levels of CTL response *in vitro*.
- Peptides 5.3 and D2 bound to HLA A\*0201 with low affinity and were variable, particularly D2.
- Substitutions in peptide D2: IlnTIaiav did not abrogate the response, but diminished it.
- In a longitudinal study, the CTL response to the variable D2 epitope diminished over time, while the response to the conserved epitope D1 stayed higher.

HXB2 Location gp160 (814–822)

**Author Location** (C consensus)

**Epitope** LLDTIAIAV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0205)

Country South Africa.

**Assay type** CD8 T-cell Elispot - IFNγ

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location gp160 (814–822)

Author Location (C consensus)

Epitope LLDTIAIAV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0205)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- LLDTIAIAV is an optimal epitope.

HXB2 Location gp160 (814–822)

Author Location gp41 (815–823 LAI)

**Epitope** LLNATDIAV

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human (A2)

References Dupuis et al. 1995

 Of two CTL clones, one reacted only with 815-823, the other with 814-823 and 815-823.

HXB2 Location gp160 (814–822)

Author Location Env (815–823)

**Epitope** LLNATAIAV

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Kmieciak et al. 1998b

 Increased CTL response to cells expressing a VV construct ΔV3 mutant compared with a full-length env gene product.

HXB2 Location gp160 (822-832)

Author Location gp41 (SF2)
Epitope VAEGTDRVIEI
Immunogen HIV-1 infection
Species (MHC) human

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of individuals that had a CTL response to this epitope (HLA presenting molecule uncertain) broken down by group: 0 group 1, 1 group 2, and 0 group 3.

**HXB2 Location** gp160 (824–832)

Author Location gp160 (828–836 WEAU)

**Epitope** EGTDRIVIEI **Epitope name** gp160 EI9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*2902, B\*4403, B\*0801

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

**Keywords** dynamics, immunodominance, acute/early infection, kinetics, characterizing CD8+ T cells,

reversion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance. WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.
- This was one of five reasonably strong responses in early infection in the patient WEAU, and the epitope sequence did not vary during the first year of the infection.

**HXB2 Location** gp160 (827–841) **Author Location** gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR Immunogen HIV-1 infection Species (MHC) human (A2) References Clerici *et al.* 1991a

 Helper and cytotoxic T cells can be stimulated by this peptide (Th4)

HXB2 Location gp160 (827–841) Author Location gp41 (834–848 IIIB) Epitope DRVIEVVQGAYRAIR

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species** (MHC) mouse  $(H-2^d, H-2^p, H-2^u, H-2^q)$ 

References Shirai et al. 1992

• In a murine system multiple class I molecules can present to CTL.

HXB2 Location gp160 (827–841) Author Location gp41 (834–848 IIIB) Epitope DRVIEVVQGAYRAIR

Immunogen vaccine

Vector/Type: vaccinia HIV component: gp160

**Species (MHC)** mouse  $(H-2^d, H-2^p, H-2^u, H-2^q)$ 

References Shirai et al. 1996b

 Multiple murine MHC can cross-present this epitope (HP53), and P18 RIQRGPGRAFVTIGK, to specific CTL.

HXB2 Location gp160 (827–841)
Author Location gp41 (834–848 IIIB)
Epitope DRVIEVVQGAYRAIR
Immunogen HIV-1 exposed seronegative

Species (MHC) human

References Pinto et al. 1995

 CTL and T helper cell reactivity in healthcare workers exposed to HIV.

**HXB2 Location** gp160 (828–836)

**Author Location** Env (829–837 subtype B)

Epitope RVIEVLQRA

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human (A\*0201)

**Keywords** binding affinity **References** Kundu *et al.* 1998a

- Ten HIV-1 + HLA A2 asymptomatic individuals were given two courses of HIV-1 MN rgp160 vaccine over a 2 year period.
- Two hundred and fifty three HIV-1 peptides of 9 or 10 aa possessing the HLA-A2.1 binding motif (Leu at position 2, Val at the C terminus) were identified in gp160, of which 25 had a high or intermediate binding affinity.
- Eleven peptides were studied that had high HLA-A2 binding affinity a CTL response was detected to 9/11 peptides in at least 1 individual.
- CTL responses after reimmunization may include recall responses individuals with vaccine cross-reactive sequences prior to vaccination showed detectable CTL responses.

**HXB2 Location** gp160 (828–836) **Author Location** gp41 (829–837 LAI)

Epitope RVIEVLQRA

**Subtype** B **Immunogen** vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

**Species** (MHC) human (A2) **References** Dupuis *et al.* 1995

• CTL from HLA-A2 positive subject react with this peptide.

HXB2 Location gp160 (828-836)

Author Location gp41 (829–837 CM243 subtype CRF01)

Epitope KVIEVAQGA
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A2)
Konwords subtype comparie

**Keywords** subtype comparisons **References** Bond *et al.* 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- 1/4 tested FSWs recognized the E clade version of this epitope, which differs from the previously defined B clade version by three amino acids, RvievLqRa.
- This epitope was only conserved in CRF01 (subtype E), and identities were rare.

**HXB2 Location** gp160 (830–854)

Author Location gp41 (831–853)

Epitope IEVVQGAYRAIIRHIPRRIRQGLERI

Immunogen HIV-1 infection

Species (MHC) human

References Price et al. 1995

• Study of cytokines released by HIV-1 specific activated CTL.

**HXB2 Location** gp160 (831–838) **Author Location** Env (830–837)

 ${\bf Epitope} \ \, {\sf EVAQRAYR}$ 

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*3303)

References Hossain et al. 2001; Takiguchi et al. 2000

- HLA-A33 a very common allele in Asia, with HLA-A\*3303 the most common among the Japanese. New A\*3303 epitopes were defined to better characterize the immune response in this population.
- The anchor motif for HLA\*3303 (A, I, L, V, F, Y in position 2 (F and Y bind most strongly), and R (K is also tolerated) in the C-terminal position) was used to define 82 potentially reactive peptides in Env; 37/82 peptides bound to A\*3303; 3/37 peptides could induce peptide-specific CTL in bulk PBMC cultures from 1/3 HLA A\*3303 positive individuals tested.
- 2/3 peptides that reacted with the bulk culture, EVAQRAYR and VIEVAQRAYR, were overlapping, with one encompassing the other, but EVAQRAYR was shown to be the one that was reactive with a CTL clone.

• CTL clones were isolated that killed target cells in a concentration dependent manner after pulsing with the EVAQRAYR peptide, that could also kill cells transfected with env expressed from a vaccinia vector. Bulk cultures were tested from six additional people, and only 2/6 reacted with this peptide, but the peptide is in a highly variable region.

**HXB2 Location** gp160 (831–838)

Author Location gp41 (320-327)

Epitope EVAQRAYR

Immunogen HIV-1 infection

Species (MHC) human (A\*3303)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** gp160 (833–841)

Author Location gp160 (837–845 WEAU)

**Epitope** VQRTCRAIL **Epitope name** gp160 VL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*2902, B\*4403, B\*0801

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

Keywords dynamics, immunodominance, acute/early in-

fection, characterizing CD8+ T cells, rever-

sion, viral fitness

References Jones et al. 2004

- Primary CD8+ T-cell response to Env, Tat and Gag and the extent, kinetics and mechanisms of viral escape were examined in three patients. Rapid escape, within weeks from infection, from HIV specific CTL responses was observed in all three patients, but the kinetics and extent of the escape differed depending on the breadth and co-dominant distribution of CTL-mediated pressure. The two patients that rapidly declined had more focused immunodominant responses, while the single patient that had low viral load and stable CD4 counts for seven years had a broad co-dominant response and less escape.
- The patient WEAU had high viral loads and rapid CD4 decline. WEAU mounted 14 detected CTL responses, with distinct patterns of immunodominance. WEAU did not control viral replication well, and escape mutations occurred early and 4/14 had changes that could have resulted in escape, and two were confirmed as escape.
- This was one of five reasonably strong responses in early infection in the patient WEAU, and the epitope sequence did not vary during the first year of the infection.

**HXB2 Location** gp160 (835–843)

Author Location Env (834–842 SF2)

Epitope RAYRAILHI

Immunogen HIV-1 infection

Species (MHC) human (B\*5101)

**Keywords** rate of progression

References Tomiyama et al. 1999

- HLA-B27, -B51, and -B57 are associated with slow progression to AIDS, while HLA-B35, -B8, -B24 are associated with a rapid progression to AIDS (Nat. Med. 2:405, 1996;Lancet 22:1187, 1986;Hum Immunol 22:73, 1988;Hum Immunol 44:156, 1995)
- 15% of Japanese populations carry HLA-B51 while HLA-B27 and -B57 are detected in less than 0.3%
- Of the 172 HIV-1 peptides with HLA-B\*5101 anchor residues, 33 bound to HLA-B\*5101, seven of these peptides were reactive with CTL from 3 B\*5101 positive individuals, and six were properly processed.
- This peptide could stimulate CTL from one person, however this CTL clone did not recognize B\*5101 positive target cells infected with HIV-1 recombinant vaccinia expressing Env, so it was not confirmed that this peptide was a properly processed epitope.

HXB2 Location gp160 (835–843) Author Location Env (835–843) Epitope RAYRAILHI Immunogen HIV-1 infection Species (MHC) human (B51)

 $\textbf{Donor MHC} \ \, \text{A03, A32, B51, B15, Cw03, Cw06, DR4,} \\$ 

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 7 epitopes from this individual that varied over time, although the internal mutations did not become fixed. The rTFrailhi variant residues arose at early time points, rIyrailhX variant residues arose at intermediate time points.

**HXB2 Location** gp160 (837–856) **Author Location** gp120 (844–863 LAI)

Epitope YRAIRHIPRRIRQGLERILL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35) References Shankar *et al.* 1996

HXB2 Location gp160 (837–856) Author Location gp41 (844–863 HXB2) Epitope YRAIRHIPRRIROGLERILL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8)

References Lieberman et al. 1992

• CTL epitope defined by T cell line and peptide mapping.

**HXB2 Location** gp160 (837–856) **Author Location** gp120 (844–863)

Epitope YRAIRHIPRRIRQGLERILL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

 HIV-specific CTL lines developed by ex vivo stimulation with peptide.

**HXB2 Location** gp160 (837–856)

**Author Location** gp120 (844–863 SF2)

Epitope YRAIRHIPRRIRQGLERILL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI gp160.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-A2, A26, B7, and B38.

HXB2 Location gp160 (842–856)

**Author Location** gp41 (SF2)

Epitope HIPRRIRQGLERALL

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2001a

- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- The only Env peptide recognized was gp41 HIPRRIRQGLER-ALL.

**HXB2 Location** gp160 (843–851)

Author Location gp41 (848–856 LAI)

Epitope IPRRIRQGL

Subtype B

**Immunogen** 

Species (MHC) human (B\*0702)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*0702 epitope.

**HXB2 Location** gp160 (843–851)

Author Location gp41 (848–856 LAI)

Epitope IPRRIRQGL

Subtype B

Immunogen

Species (MHC) human (B7)

Keywords mother-to-infant transmission

References Brander & Walker 1995

• Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.

HXB2 Location gp160 (843–851)

**Author Location** 

Epitope IPRRIRQGL Immunogen HIV-1 infection Species (MHC) human (B7)

**Keywords** immunodominance, escape **References** Soudeyns *et al.* 1999

- Following primary infection, progressive diversification and accumulation of mutations of HIV-env nucleotide sequences was observed, focused in V2 in one individual and in V8 in another.
- The patient with the V2 diversification showed only transient CTL against Env and Nef.
- The patient with the V8 diversification had an immunodominant CTL response to V8 epitope IPRRIRQGL, and multiple escape variants emerged within a year: ipTrirqgl and ipTrirqgF, which abrogated the CTL response *in vitro*, and also iprrLqgl and iprrirqDl which gave diminished responses.

**HXB2 Location** gp160 (843–851) **Author Location** gp41 (848–856 LAI)

Epitope IPRRIRQGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

 $\textbf{Keywords} \ \ \text{subtype comparisons}$ 

**References** Cao *et al.* 1997a

- The consensus peptide of clades A, B, D, and F is IPRRIRQGL.
- The consensus peptide of clade C is iprrirqgF, and it is equally reactive.

HXB2 Location gp160 (843–851)

Author Location gp41 (848–856 subtype B)

Epitope IPRRIRQGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Keywords** subtype comparisons, acute/early infection **References** Wilson *et al.* 1998b

- The extent of CTL interclade cross-reactivity from CTL isolated from individuals newly infected with B clade virus was studied, and extensive cross-reactivity was observed.
- Two HLA B7 individuals had CTL response to B\_LAI, A\_92UG037 and C\_92BR025 gp160, but were B clade strain MN non-responders – the authors note that the B7 epitope IPRRIRQGL is conserved between the LAI and clade A and C strains, but that MN has a non-conservative Arg to Thr substitution at position three that may be contributing to the specificity of the response in the HLA B7 individuals.

**HXB2 Location** gp160 (843–851)

Author Location gp41 (843–851 HXB2)

Epitope IPRRIRQGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords rate of progression, immunodominance

References Hay et al. 1999

- CTL response to IPRRIRQGL was the immunodominant response in a rapid progressor there was a subdominant response to SPAIFQSSM in Pol, and interestingly, no response to commonly immunodominant HLA A\*0201 epitope SLYNT-VATL, although this individual was HLA A\*0201.
- The individual showed a strong initial CTL response at the time of the initial drop in viremia, but it was quickly lost, although memory cells persisted.

- Despite the initial narrow response to two epitopes, no other CTL responses developed.
- No HIV-specific lymphoproliferative responses were detected in this patient, and neutralizing antibody response was weak.
- Variants were observed *in vivo*, the most common form of the viral epitope at presentation at 3 months was the only form that did not elicit a CTL response: iprrTrqgl; the other forms detected were iprrirqgF, iprriLqgF, VprrirqgF and they could elicit a CTL response although the response to iprriLqgF was reduced.
- A second rapid progressor had a detectable CTL response exclusively to this epitope.

**HXB2 Location** gp160 (843–851)

Author Location gp41 (subtype A)

Epitope IPRRIRQGF

Subtype A

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Keywords** subtype comparisons

References Cao et al. 2000

- HIV-1 subtypes A and D dominate the Ugandan epidemic, and a vaccine trial using B clade antigen is underway – this study addresses relative levels of cross-reactive CTL responses in Ugandans to A, D, and B clade recombinant vaccinia viruses expressing Gag, Env, Pol, RT or Nef from HIV-1 clades A, B, and D.
- Proteins corresponding to the subtype of the infecting strains tended to trigger higher levels of CTL response measured by percent specific lysis, but there was extensive inter-subtype cross-reactivity with B clade proteins and the co-circulating subtype.
- This optimal epitope sequence, recognized by CTL derived from a Ugandan with an A subtype infection, is cross-reactive with subtypes A and B, but not in subtype D.

**HXB2 Location** gp160 (843–851)

**Author Location** gp41

Epitope IPRRIRQGL

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Keywords** rate of progression, acute/early infection

References Islam et al. 2001

- Subject 053i was followed longitudinally from acute infection through death, and had rapid progression to AIDS.
- This individual had a dominant response to IPRRIRQGL with strong *in vivo* activated responses and *in vitro* stimulated memory responses and a subdominant response to SPAIFQSSM during the course of disease progression (4 Years), the functional CTL responses were lost and no sequence variation occurred with in both epitopes.
- At 3 months post-presentation, seven IPRRIRQGL CTL clones were obtained, five used the T-cell receptor V $\beta$  6S1 and J $\beta$  2.7 and had the CDR3 WAASS, two used V $\beta$ 16S1, ERSPPGD, J $\beta$  2.7 and one CTL clone isolated at 39 months was V $\beta$  14S1, CR3 PTAAG, and J $\beta$  2.1 all of these clones persisted over the course of the infection, even to time of death, despite the loss of CTL functional responses over time.

**HXB2 Location** gp160 (843–851)

Author Location gp41 (843–851 SF2)

Epitope IPRRIRQGL Immunogen HIV-1 infection Species (MHC) human (B7)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B7+ individuals that had a CTL response to this epitope broken down by group: 2/4 group 1, 1/3 group 2, and 1/1 group 3.

**HXB2 Location** gp160 (843–851)

Author Location gp41 (848–856)

Epitope IPRRIRQGL

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (B7)

**Keywords** HIV exposed persistently seronegative (HEPS), immunodominance

References Kaul et al. 2001a

- IPRRIRQGL cross-reacts with clades A, B and D.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B7 women, 2/5 HEPS and 5/6 HIV-1 infected women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in 2 of the 5/6 HIV-1 infected women that responded to the epitope, but in neither of the 2/5 HEPS cases.
- Subject ML 1203 started with CTL responses to A\*6802 DTVLEDINL and to B7 FPVTPQVPLR prior to seroconversion, and upon seroconversion acquired additional responses to A\*6802 ETAYFILKL which became dominant, B7 TPGPGV/IRYPL, B7 IPRRIRQGL, and B7 SPRTLNAWV.

HXB2 Location gp160 (843–851) Author Location gp41 (843–851) Epitope IPRRIRQGL Immunogen HIV-1 infection Species (MHC) human (B7)

**Keywords** rate of progression, acute/early infection **References** Day *et al.* 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

HXB2 Location gp160 (843–851)

Author Location gp41 (SF2)

Epitope IPRRIRQGL

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Altfeld et al. 2000

• This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.

**HXB2 Location** gp160 (843–851)

Author Location gp41 (842–852)

Epitope IPRRIRQGL

Epitope name B7-IL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), immunodominance, acute/early infec-

tion

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- Only two epitopes were detected during acute infection in patient AC-06, B7 restricted gp41 epitope IPRRIRQGL and Gag GPGHKARVL. GPGHKARVL was the first targeted peptide, and remained immunodominant through the 34 month study period.

• 6/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection - 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 2/4 individuals had detectable responses to this epitope after STI.

HXB2 Location gp160 (843-851)

Author Location gp41

Epitope IPRRIRQGL

Epitope name B7-IL9(gp41)

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7) **Donor MHC** A24, B7, B27

Keywords HAART, ART, supervised treatment interrup-

tions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hivweb.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples of the 15 patients, and an additional 8 responses were detected only in LN. The total magnitude of the response was similar in LN and PB, but the percentage of CD8+ T cells in the LN is lower so the number of HIV-specific cells per million CD8+ T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- · Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- · Breakdowns of epitope responses were shown for 4 individuals. Patient C displayed the greatest response to B27-KK10(p24), and in decreasing order also responded to A24-RW8(Nef), B7-IL9(gp41), A24-RL9(gp41), A24-YL8(gp41), and B7-TM9(Nef).

HXB2 Location gp160 (843-851)

**Author Location** Env

Epitope IPRRIRQGL

Epitope name EW10

Immunogen HIV-1 infection

Species (MHC) human (B7)

Assay type Chromium-release assay, Flow cytometric T- Author Location gp41

cell cytokine assay

**Keywords** class I down-regulation by Nef

References Bobbitt et al. 2003

• Nef, through Nef-mediated MHC-1 down-regulation, is not the only viral protein to influence levels of HIV-specific CTL recognition. The Rev L60F mutation, a common natural variant, can decrease CTL recognition of late gene products. The Rev mutation impacts the early to late gene switch, reducing late gene product production (Gag, Pol, Env, Vpu, Vpr and Vif), while increasing Nef production, both of which reduce the

impact CTL recognition of late gene products. As expected, Rev L60F rendered HIV infected cells more resistant to CTL that recognized epitopes from the late proteins Env and Gag. Gag expression is reduced more than Env, and Gag-specific CTL were more profoundly affected. Conversely CTL against an epitope in an early gene product, Tat, were more effeciently recognized when infected with viruses carrying the Rev L60F mutation.

Patients in the asymptomic phase with active immune responses had more CTL resistant viruses, with lower Rev activity, lower Gag expression and greater resistance to Gag-specific CTL killing, while viruses isolated from people with AIDS were more sensitive to CTL killing.

HXB2 Location gp160 (843–851)

Author Location gp41 (843–851)

Epitope IPRRIRQGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A1, A3, B7, B14, Cw\*0702, Cw\*0802

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords binding affinity, acute/early infection, earlyexpressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

HXB2 Location gp160 (843–851)

Epitope IPRRIRQGL

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** HIV exposed persistently seronegative

(HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- 3/9 HLA B7+ infection-resistant men, compared to 0/4 preseroconversion men who went on to become infected, reacted to this epitope.

HXB2 Location gp160 (843–851)
Author Location Env (333–341)
Epitope IPRRRIRQGL
Immunogen HIV-1 infection
Species (MHC) human (B7)
Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 3/7 patients recognized this epitope.

HXB2 Location gp160 (843–851)
Author Location (B consensus)
Epitope IPRRIRQGL
Epitope name IL9
Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

**Donor MHC** A31, A68, B07, B70, Cw7, Cw1

Assay type Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

HXB2 Location gp160 (843–851)
Author Location gp41
Epitope IPRRIRQGL
Subtype B
Immunogen HIV-1 infection

Species (MHC) human (B7)
Country United Kingdom.

**Assay type** Tetramer binding, T-cell Elispot, Intracellular cytokine staining

**Keywords** rate of progression, acute/early infection, characterizing CD8+ T cells, immune dysfunction

**References** Papagno *et al.* 2004 Acute HIV-1 infection induces massi

Acute HIV-1 infection induces massive activation of HIV-specific and non-HIV-specific CD8+ T-cells resulting in differentiation of these cells. High differentiation of CD8+ T-cells is correlated with disease progression. Differentiation is a natural process but it can be driven by elevated immune activation, such as in HIV infection.

Author Location gp160 (843–851)
Author Location gp41 (333–334)
Epitope IPRRIRQGL
Epitope name IPR
Immunogen HIV-1 infection
Species (MHC) human (B7)

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401, Cw\*0702, DR17, DR15, DR51, DR52, DQ2, DQ6

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** rate of progression, escape **References** Oxenius *et al.* 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant epitopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This was one of 8 reactive epitopes found not to vary over time.

HXB2 Location gp160 (843–851) Author Location gp120 Epitope IPRRIRQGL

Epitope name IL9 Immunogen

Species (MHC) (B7)

**Keywords** review, immunodominance, escape, acute/early infection, early-expressed proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

This review discusses the importance of 3 factors that impact
the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA
class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides
listed as immunodominant in acute HIV-1 infection.

HXB2 Location gp160 (843–851) Author Location gp160 (clade A, B, C, D) Epitope IPRRIRQGL **Subtype** A, B, C, D **Immunogen** HIV-1 infection

Species (MHC) human

**Donor MHC** A\*6802, A\*6802, B\*1303, B\*1401, Cw\*0602, Cw\*1701

Country Kenya.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Other

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References McKinnon et al. 2005

- Interclade cross-reactivity of clades A, B, C, and D was tested using recombinant vaccinia-based IFN-gamma Elispot assay. 47/74 women had a positive Elispot response to at least one clade, and cross-clade responses were frequent. Clade A responses were most frequent, as expected, as clade A dominates the Kenyan epidemic (85% of the 47 women responded to clade A Env, 74% to B, 62% to C, and 43% to D). Cross-clade CD8 T-cell responses were common and directed at conserved epitopes.
- There was a greater magnitude of response to A clade peptides in individuals who responded to more than 1 clade; a 2-fold higher response was observed in clade A in 36% (9/25) of these individuals, and the response to A peptides was never lower. IPRRIRQGL responses were detected in all 4 clades in 1 woman: clade A gave a high response; IPRRIRQGL was identical clade B; clade C and D had lower responses and carried variant peptides IPRRIRQGf and IvRRIRQGL.

**HXB2 Location** gp160 (843–851)

**Author Location** gp160

Epitope IPRRIRQGL Subtype A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: vaccinia Strain: A clade, B clade, D clade NDK, C clade consensus HIV component: Env

Species (MHC) human

**Donor MHC** A\*0205, A\*3402, B\*4201, B\*5802, Cw\*0602, Cw\*1701

Country Kenya.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Other

**Keywords** subtype comparisons, variant cross-recognition or cross-neutralization

References McKinnon et al. 2005

- Interclade cross-reactivity of clades A, B, C, and D was tested using recombinant vaccinia-based IFN-gamma Elispot assay. 47/74 women had a positive Elispot response to at least one clade, and cross-clade responses were frequent. Clade A responses were most frequent, as expected, as clade A dominates the Kenyan epidemic (85% of the 47 women responded to clade A Env, 74% to B, 62% to C, and 43% to D). Cross-clade CD8 T-cell responses were common and directed at conserved epitopes.
- There was a greater magnitude of response to A clade peptides in individuals who responded to more than 1 clade; a 2-fold higher response was observed in clade A in 36% (9/25) of these individuals, and the response to A peptides was never lower. An IPRRIRQGL response was detected in a women who had Env responses to 3 clades, A, B, and C, and clade A and B gave

the highest responses. IPRRIRQGL was identical in A and B, had the form IPRRIRQGf in C, and IvRRIRQGL in D.

**HXB2 Location** gp160 (845–856)

Author Location gp41 (852–863 HXB2)

Epitope RRIRQGLERILL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A30, B8)

References Lieberman et al. 1992

• CTL epitope defined by T cell line and peptide mapping.

**HXB2 Location** gp160 (845–856)

Author Location gp41 (852-863 LAI)

Epitope RRIRQGLERILL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Shankar et al. 1996

HXB2 Location gp160 (846-854)

**Author Location** 

Epitope RIRQGLERA

**Epitope name** Env-RA9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0205)

**Donor MHC** A\*0205 A\*3002 B\*1402 B\*5301 Cw\*0401 Cw\*0802

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- This epitope was newly defined in this study.
- Patient 00RCH28 was African American, not on HAART, had a viral load of 5900 and CD4 count of 889, and she also recognized IN(219-227), KIQNFRVYY, A\*3002.
- Among HIV+ individuals who carried HLA A02, 6/21 (29%) recognized this epitope.

**HXB2 Location** gp160 (846–854)

Author Location gp41 (335–343)

Epitope RIRQGLERA

Immunogen HIV-1 infection

Species (MHC) human (A\*0205)

**Keywords** optimal epitope

**References** Frahm *et al.* 2007

**HXB2 Location** gp160 (848–856)

**Author Location** gp160 (848–856) **Epitope** RQGLERALL

Immunogen

Species (MHC) human (B8)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B8 epitope.

HXB2 Location gp160 (849-856) Author Location gp41 (849–856) Epitope QGLERALL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8)

**Donor MHC** A1, A1, B8, B14, Cw7, Cw8 Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, earlyexpressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFNsecreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- · All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

## II-B-22 Env CTL/CD8 + epitopes

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Immunogen computer prediction **Species (MHC)** (A\*0201, B\*3501)

> **Keywords** subtype comparisons, computational epitope prediction

References Schönbach et al. 2002

• Computational methods (artificial neural networks, hidden Markov models, binding matrices based on HLA association rates) were used to identify HLA-A\*0201 and HLA-B\*3501 HIV T-cell epitope candidates from 533 Gag, Env and Pol sequences of which 374 were derived from HIV-1, 97 were derived from HIV-2 and 62 from SIV. Comparisons to known epitopes and between clades were made.

**HXB2 Location** Env **Author Location Epitope** 

Subtype A, B, C Immunogen vaccine

*Vector/Type:* canarypox prime with gp120 boost, canarypox prime with gp160 boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Gag, gp120, gp41, Nef, Pol

Species (MHC) human (A1, A2, A24, B62, A25, A26, A30, A31, B8, B17, B39, B51, B57, B60, B70)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Ferrari et al. 2001

- Different HIV strains were used for different regions: gp41 LAI, Gag LAI, gp120 MN, gp120 SF2
- HLA-B62 responses dominated the responses against an Env vaccine in an individual (022JAV) who was HLA A2, A26, B35, B62. The strongest response was against the MN peptide 381-400; a response diminished by half was observed against vaccinia expressed clade A and clade C relative to clade B.
- Class I presentation of Env CTL responses in vaccinee 022A12K: A25 > B39, A1 and B8 were undetectable.
- Class I presentation of Env CTL responses in vaccinee 022A12N: B57 » A2 > A26 and B60.
- Class I presentation of Env CTL responses in vaccinee 034GP3: A31 > A24 > B62 > B51.
- Class I presentation of Env CTL responses in vaccinee 0348PP: B17 > B70, A1 and A30 were undetectable.

**HXB2 Location** Env

**Author Location** gp120 (303–327)

**Epitope** 

Immunogen HIV-1 infection

**Species (MHC)** human (A2, A3, A11, B27)

**Keywords** subtype comparisons References Ferrari et al. 2000

- One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.
- For this cluster of epitopes spanning the tip of the V3 loop, they suggest including a sequence from each clade.

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype B

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost, canarypox prime with gp160 boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Gag, gp120, gp41, Nef, Pol

Species (MHC) human (A2, B8)

Keywords vaccine-induced epitopes

References Ferrari et al. 2001

- Different HIV strains were used for different regions: gp41 LAI, Gag LAI, gp120 MN, gp120 SF2
- No HLA-A\*0201 or B8 responses were made against the Env vaccine in individuals carrying these alleles, despite these being common presenting molecules for CTL responses to natural infections.

**HXB2 Location** Env Author Location Env

> **Epitope** Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*35)

**Keywords** rate of progression

References Jin et al. 2002

- Patients with HLA-B\*35 variants B\*3502, B\*3503, B\*3504, and B\*5301 tend to proceed to AIDS more quickly than those with B\*3501.
- Of 32 patients with HLA-B\*35 alleles CD8+ CTL responses were quantified using an intracellular cytokine staining assay – 75% had responses to Pol, 69% to Gag, 50% to Nef, and 41%
- The overall magnitude of CTL responses did not differ between those bearing B\*3501 and the others. A higher percentage of Gag responses was observed in those that had lower RNA levels that carried B\*3501, and there was a negative association with viral load and CTL activity. The data is consistent with higher levels of CTL responses contributing to protection in B\*3501 individuals, but not in B\*3502, B\*3503, B\*3504, and B\*5301 individuals.

**HXB2 Location** Env

Author Location gp41 (842–850 IIIB, BH8)

**Epitope** 

Immunogen HIV-1 infection Species (MHC) human (B7)

References Pantaleo et al. 1997; Soudeyns & Pantaleo

• Clonotype-specific PCR and analysis of in vivo HIV-specific CTL showed that in early infection HIV-specific CTL clones preferentially accumulate in blood rather than lymph nodes and that they accumulate prior to down-regulation of virus.

**HXB2 Location** Env

Author Location gp120 (318-327 IIIB)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB

HIV component: gp120

Species (MHC) mouse (H-2 Dd)

Assav type Cytokine production, Intracellular cytokine staining, Th support of CTL response,

Chromium-release assay

Keywords epitope processing, Th1, vaccine antigen de-

sign

References Vatakis et al. 2005

• Mice were vaccinated with three DNA epitope vaccines, differing in the affinity of the helper epitope to the MHC class II molecule. It was observed that a TH epitope with lower affinity decreased the magnitude of the CTL responses and decreased the numbers of epitope-specific T-helper cells and CTLs. Also, cytokine secretion and proliferative responses were diminished.

**HXB2 Location** Env

Author Location gp160 (MN)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade MN HIV component: gp120, gp160

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Vinner et al. 1999

- Mammalian codon optimization renders gp160 expression Rev independent, increases gp160 expression levels, and DNA vaccination of BALB/c mice yields a higher antibody response with an earlier onset than wild type.
- Secreted gp120 gave higher antibody titers than membrane bound gp160.
- In contrast to antibodies, synthetic codon-optimized DNA did not alter the CTL response, wild type genes generated equally strong CTL responses.

**HXB2 Location** Env

**Author Location** (IIIB)

**Epitope** 

Immunogen vaccine

Vector/Type: peptide HIV component: V3 Adjuvant: Cholera toxin (CT), GM-CSF, IL-4

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Kato et al. 2000

- A multicomponent peptide vaccine VC1 with cholera toxin adjuvant was given to mice.
- Immunization of BALB/c mice with VC1 and CT induced a strong CTL response which was enhanced by IL-12 expressing plasmids.
- Immunization with VC1 and CT resulted in HIV-1 specific IgA antibody responses, which were increased by the combination of IL-4 or GM-CSF expressing plasmids.

**HXB2 Location** Env

Author Location gp160 (IIIB)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB HIV component: gp160 Adjuvant: PLG

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Kaneko et al. 2000

- A PLG-microparticle encapsulated DNA encoding gp160 was given to mice.
- Oral DNA vaccination of BALB/c mice induced mucosal and systemic gp160 glycoprotein-specific cellular and humoral immune responses, and mice vaccinated orally had higher resistance to HIV-env expressing vaccinia intrarectal challenge than mice vaccinated i.m.

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Immunogen vaccine

Vector/Type: DNA with CMV promotor with cationic liposome HIV component: gp160, Rev

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Ishii et al. 1997

• pCMV160/Rev is a DNA vaccine candidate carrying gp160 and Rev linked to a cytomegalovirus (CMV promotor)

**HXB2 Location** Env

Author Location Env Epitope

Immunogen vaccine

Vector/Type: adeno-associated virus (AAV) HIV component: Env, Rev, Tat Adjuvant: IL-2

**Species (MHC)** mouse (H-2<sup>d</sup>) **References** Xin *et al.* 2001

- An AAV vector expressing HIV-1 env, tat, and rev genes (AAV-HIV vector) was used to vaccinate BALB/c mice.
- A single injection stimulated and long lasting serum IgG, fecal IgA, and HIV-specific CTL.
- Boosting enhanced the humoral response, and IL2 enhanced T-cell immunity.

HXB2 Location Env Author Location Env Epitope

Immunogen vaccine

Vector/Type: vaccinia, influenza Strain: B clade IIIB HIV component: Env, V3

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Gonzalo et al. 1999

The use of two different live vectors for priming and boosting
has a synergistic effect on the immune response against HIV-1
a 5-6 fold enhanced CTL response in Balb/c mice occurred
when they were immunized with rec influenza virus (Flu-Env)
expressing the V3 loop epitope from HIV-1 strain IIIB, and
boosted with a vaccinia virus recombinant (VV-Env) expressing the complete HIV-1-IIIB env protein, comared to either
immunogen alone.

**HXB2 Location** Env

**Author Location** Env (subtype B)

Epitope Subtype B Immunogen vaccine

Vector/Type: rabies virus Strain: B clade 89.6, B clade NL43 HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>)

References McGettigan et al. 2001

- BALB/c were immunized with a replication competent recombinant rabies virus (RV) vaccine expressing HIV-1 gp160.
- A single vaccination induced induced strong and long-lasting (4.5 months) gp160-specific CTL cytotoxic responses.
- Although the greatest specific lysis was achieved when the
  vaccine strain was also used as the *in vitro* the target strain to
  assess the response, there was extensive CTL cross-reactivity
  against other B clade HIV-1 envelope proteins, implying CTL
  recognition of multiple epitopes within the HIV-1 envelope
  protein.

HXB2 Location Env Author Location gp120 (V3) Epitope Subtype B

Immunogen vaccine

Vector/Type: DNA, polyepitope Strain: B clade BRVA, B clade IIIB, B clade JY1, B clade LR150, B clade MN, B clade RF HIV component: V3

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type CD8 T-cell Elispot - IFN $\gamma$ References Vázquez-Blomquist *et al.* 2003

- Priming mice with recombinant MVA and boosting with fowlpox was shown to increase the number of specific IFN-gamma secreting cells relative to reversing the order (fowlpox prime, MVA boost) or priming with a Semliki Forest Virus DNA vector and boosting with recombinant MVA or fowlpox. The authors speculate why the order might be important. Fowlpox has more proteins, so there may be more CTL epitope competition; alternatively pox viruses may modulate the immune response through chemokine homologs.
- The antigen tested was a V3 loop polyepitope vaccine combining multiple V3 loop variants given by an intraperitoneal route to BALB/c mice.

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: gp120 Adjuvant: Cholera toxin (CT)

**Species (MHC)** mouse (H-2D<sup>d</sup>)

Assay type T-cell Elispot, Chromium-release assay

References Bagley et al. 2003

BALBc mice were immunized intramuscularly with single plasmids encoding gp120, or cholera toxin catalytic domain (CTA1) and gp120, or with a dicistronic DNA vaccine expressing both CTA1 and gp120. Vaccination including CTA elicited stronger and longer lasting Ab responses and T-cell responses to gp120.

**HXB2 Location** Env

**Author Location** Env (SIV)

**Epitope** 

Immunogen SIV infection

Species (MHC) macaque (Mamu-A\*11, Mamu-B\*03, Mamu-B\*04, Mamu-B\*17)

References Dzuris et al. 2000

• Cell binding assays for Mamu molecules were employed to describe the peptide binding motifs for Mamu-A\*11, -B\*03, -B\*03, -B\*04, and -B\*17 CTL epitopes – a similarity for Mamu-A\*11 and -B\*03 and human HLA-B\*44 and -B\*27, respectively, was observed – all epitopes studied were SIV epitopes, so not specifically listed here.

**HXB2 Location** Env

Author Location gp160 (LAI, MN)

**Epitope** 

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Gag, gp120, gp41, Protease

Species (MHC) human

References Belshe et al. 1998

• The live canarypox vaccine ALVAC-HIV(vCP205) carrying • The vaccine used was a rec canarypox with HIV-1 gp120 MN, MN gp120, LAI gp41, Gag and Protease, and boosted with SF-2 rpg120, was given to HIV-1 seronegative volunteers -HIV-specific Env or Gag CD8+ CTL were detected in 64% of the volunteers.

**HXB2 Location** Env

Author Location gp160 (LAV)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords epitope processing, dendritic cells

References Zheng et al. 1999

- Protein delivery (gp160 LAV, p66 LAV, and p24 NY5) to human dendritic cells (DC) with liposomes provides enhanced memory CTL response relative to delivery of protein alone.
- · Chloroquine administration enhanced epitope presentation, and brefeldin A and peptide aldehyde inhibitors inhibited antigen presentation, suggesting epitopes were processed by classical proteasome pathway.

**HXB2 Location** Env

**Author Location** Env (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression, Th1

References Wasik et al. 2000

- HIV+ infants that progressed rapidly to AIDS had lower Th1 responses and decreased production of IL-2, as well as betachemokines, relative to other HIV+ infants.
- No HIV+ infants had no demonstrable CTL at birth, but Th1 responses accompanied by CTL responses developed in children with slowly progressive disease, and not in rapid progressors.
- CTLp frequencies were determined by limiting dilution using autologous B cells infected with vaccina/HIV constructs.

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Soudeyns et al. 2000

- Analysis of T cell receptor beta chain variable region repertoire indicates that antiretroviral therapy (ART) and highly active antiretroviral therapy (HAART) decrease global CD8 T cell oligoclonality during primary HIV infection.
- A sharp decline in HIV-1 gp120-specific CTL clones was observed in HAART-treated subjects.

**HXB2 Location** Env

**Author Location** Env (LAI, MN)

**Epitope** 

Immunogen vaccine

Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag,

gp41, Protease, V3

Species (MHC) human

References Salmon-Ceron et al. 1999

- tm/gag/protease LAI (vCP205), alone or with p24E-V3 MN synthetic peptide (CLTB-36))
- Twenty HIV negative subjects were vaccinated in phase I trial with combinations of vCP205 and CLTB-36.
- Immunization with vCP205 induced HIV-1-specific ABs to gp120, V3, and p24 antigens, and CTL immune responses against vCP205 were detected after the fourth immunization in 33% of the subjects against Env, Gag and Pol, but the CLTB-36 peptide did not produce AB or CTL immune responses against p24 or gp160.

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords TCR usage

References Gamberg et al. 1999

- 13/13 subjects with advanced HIV infections showed CD8 T cell proliferation and differentiation of CTL in vitro, and six individuals showed HIV-specific responses to Gag, Pol, Env or Nef antigens.
- Data suggests that the functional and genetic integrity of the CD8 T cell repertoire (TCR betaV gene intrafamily genetic diversity) remains intact through advanced HIV infection, although HIV-specific CTL activity decreases.

**HXB2 Location** Env

Author Location Env (LAI, MN)

**Epitope** 

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade LAI, B clade SF2 HIV component: Env, Gag, Nef, Protease

Species (MHC) human

References Gorse et al. 1999b

- The vaccine used was rec canarypox expressing HIV-1 env, gag, pol, nef and protease (vCP300) with or without administration of HIV-1 SF-2 rgp120.
- In vitro inducible CTL activity against HIV-1 Env, Gag, Pol, and Nef antigens was observed in 79% (15 of 19) of vaccine recipients.
- The combination of vCP300 and vP1291 together resulted in an overall increase in CTL induction and detection sensitivity.

**HXB2 Location** Env

**Author Location** Env (LAI)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons References Buseyne et al. 1998b

• In infants with positive CTL responses, most responses showed cross-clade reactivity with somewhat diminished recognition of epitopes from different subtypes.

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB

HIV component: gp120, gp160

Species (MHC) macaque

References Shiver et al. 1997

 DNA vaccinations of Rhesus monkeys with a gp120 or gp160 DNA vaccine elicited a strong CD8 cytotoxic T cell response.

HXB2 Location Env

**Author Location** gp160

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) macaque

References Kent et al. 1997b

- Macaques can be infected with HIV, and clear the infection within 6 months, so it is of interest to examine their initial immune response.
- A strong CTL response against env, pol and gag antigens can be detected.
- The CTL response peaked by 4 weeks and declined dramatically by 8 weeks.
- The response in the lymph nodes and peripheral blood was comparable.

HXB2 Location Env

**Author Location** gp160

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env,

Gag, Pol, Vif Adjuvant: B7, IL-12

Species (MHC) mouse

References Kim et al. 1997c

- A gag/pol, vif or env DNA vaccine, when delivered in conjunction with the plasmid encoding the co-stimulatory molecules B7 and IL-12, gave a dramatic increase in both the cytotoxic and proliferative responses in mice.
- When IL-12 was present, CTL response could be detected even without *in vitro* stimulation.

**HXB2 Location** Env

Author Location gp160

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env,

Gag, Pol, Vif Adjuvant: B7, IL-12

Species (MHC) mouse

References Kim et al. 1997d

- A gag/pol or env DNA vaccine, when delivered in conjunction with the plasmid encoding the co-stimulatory molecules CD86, gave a dramatic increase in both the cytotoxic and proliferative responses in mice.
- When CD86 was present, CTL response could be detected even without *in vitro* stimulation.

**HXB2 Location** Env

Author Location gp120 (HXBc2)

Epitope

Immunogen vaccine

Vector/Type: DNA prime with gp160 boost Strain: B clade HXBc2 HIV component:

gp160

Species (MHC) macaque

References Letvin et al. 1997

- Vaccination of Macaques mulatta (Rhesus monkeys) with an HXBc2 env DNA prime and a protein boost elicited a T cell proliferative response, a CTL response, and type-specific neutralizing antibodies.
- Vaccinated animals challenged with SHIV-HXB2 were protected from infection.

**HXB2 Location** Env

Author Location gp120 (MN)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade MN

HIV component: Env, Rev

Species (MHC) human

References MacGregor et al. 1998

- An HIV DNA env and rev vaccine given to 15 asymptomatic HIV+ individuals at three different dosages, 30, 100 or 300 ug, was safe.
- The CTL response to gp120 was enhanced in 0/4 patients in the 30 μg group, 2/3 patients in the 100 μg group, and 0/3 in the 300 μg group – but the non-responding patients in the 300 μg group had a strong CTL response prior to vaccination, and the CTL results are inconclusive.

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Trickett et al. 1998

- Twelve HIV-1 infected patients were re-infused with their own lymphocytes, cryopreserved from an earlier time point in the infection.
- Improvement in CD4+ and CD8+ T cells was seen in 7/12, and an increase in the CTL response to Env was seen in one patient.

**HXB2 Location** Env

Author Location gp120 (LAI)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Legrand et al. 1997

- Seventeen recently infected patients were tested for CTL response to HIV proteins Env, Gag, Pol, Rev, Nef, Vif and Tat.
- An early response (within a month following PI) was noted in 87% of the subjects to Gag, 75% to Env, and 50% to Nef.
- Early responses to Pol, Rev, Vif and Tat were rare.

**HXB2 Location** Env

Author Location gp120 (LAI)

Epitope Subtype B

Immunogen vaccine

Strain: B clade LAI, B clade MN, B clade SF2 HIV component: gp160

Species (MHC) human

References Corey et al. 1998

- Vaccinia-naive subjects were vaccinated with vaccinia-gp160 LAI and boosted with gp120 SF2, LAI, MN, or 160 MN.
- 26/51 had an anti-Env CTL response, and those that were boosted with gp120 tended to produce Abs that neutralized autologous laboratory strains with some cross-reactivity.

**HXB2 Location** Env

**Author Location** Env (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons References Betts et al. 1997

- 6/8 individuals from Zambia infected with C clade virus had CTL that were able to make response to B clade HIV-1 IIIB vaccinia-expressed Gag, Pol and Env proteins.
- A vigorous cross-clade response was not limited to a particular protein, and the level of recognition of different proteins varied among the six patients.

**HXB2 Location** Env **Author Location** Env

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References De Maria et al. 1997

- CD3+ cells that also carry a natural killer cell receptor (NKR+) can exhibit down regulation of T cell function.
- · Anti-NKR IgM MAb masked this inhibitory function and increased HIV-1 specific CTL activity in phytohemagglutininactivated PBMC cultured in the presence of IL-2 from 3/5 patients, and in one other case anti-NKR MAb brought HIV-1 specific CTL activity to detectable levels.

**HXB2 Location** Env

**Author Location** Env (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression References Betts et al. 1999

• This study demonstrated an inverse correlation between HIV Type I plasma viral load and CTL activity directed against HIV-1 Pol, and stronger combined effects of Pol- and Envspecific CTL, in long-term survivors (LTS) of HIV-1 infection.

**HXB2 Location** Env

**Author Location** Env (LAI)

**Epitope** Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Buseyne et al. 1998a

Vector/Type: vaccinia prime with gp120 boost • This study showed a correlation between strong CTL memory and breadth of response in 7-12 month old infants, and remaining AIDS-free for the first year of life, higher absolute CD4 and CD8 cells, and lower viral load.

**HXB2 Location** Env

Author Location Env

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

References Goh et al. 1999

- 13/37 exposed uninfected individuals with repeated high-risk sexual exposure had HIV-1 specific CTL against Env, Gag, Pol, or a combination of proteins - CTL activity was correlated with a CCR5 wildtype genotype.
- In this group, the highest CTLp frequencies were directed at Gag, but the most common response was to Env and four individuals had responses to multiple HIV-1 proteins.

**HXB2 Location** Env

**Author Location** Env (LAI, MN)

**Epitope** 

Immunogen vaccine

*Vector/Type:* canarypox *HIV component:* Gag, gp120, gp41, Nef, Protease, RT

Species (MHC) human

References Evans et al. 1999

• A Canarypox vaccine expressing gp120, gp41, Gag, Protease, Nef and Pol CTL epitopes gave rise to CTL that could be detected in 61% of the volunteers – responses to Gag, Env, Nef and Pol were detected 3-6 months after the last vaccination.

HXB2 Location Env

Author Location Env (LAI)

**Epitope** Subtype B

Immunogen vaccine

Vector/Type: DNA prime with vaccinia boost Strain: B clade LAI HIV component: Env,

Gag

Species (MHC) macaque

Keywords Th1, Th2

References Kent et al. 1998

- · Priming with an HIV-DNA vaccine and boosting with a vaccinia construct induced greater levels of HIV T cell immunity than either vaccine alone.
- The proliferative response to Env and Gag after the DNA vaccination had a mean SI of 1.5-4, but after boosting with rHIVfowlpox virus, there was a 6-17 fold increase in the mean SI for HIV Gag and Env. The T help response happened despite a fall in antibody titers, suggesting that the Th response was primarily Th1, not Th2. The CTL response was also enhanced.

**HXB2 Location** Env

Author Location Env (LAI, MN)

**Epitope** 

Immunogen vaccine

Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag, gp120, gp41, Protease

Species (MHC) human

References Salmon-Ceron et al. 1999

 A live attenuated canarypox vector expressing MN gp120 and LAI gp41/gag/protease could induce CTL and a lymphoproliferative response in healthy, uninfected volunteers.

**HXB2 Location** Env

**Author Location** Env (MN)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env, Gag, Pol Adjuvant: CD80, CD86

Species (MHC) chimpanzee

References Kim et al. 1998

• The study explores the use of co-stimulatory molecules coexpressed with an HIV-1 immunogen in a DNA vaccine to enhance the immune response – co-expression of CD86, but not CD80, dramatically increased both HIV Env and Gag/Pol specific CTL and Th proliferative responses.

**HXB2 Location** Env

**Author Location** gp120 (IIIB)

Epitope

Immunogen vaccine

Vector/Type: Semliki-Forest Virus with viruslike particle boost Strain: B clade IIIB HIV component: Gag, gp120

Species (MHC) macaque

References Notka et al. 1999

- Immunization of SIV Pr56Gag-derived VLPs with HIV-1 gp120 anchored on their surface induced Abs, CTL and Th responses to HIV gp120; priming with the HIV antigens in Semliki-Forest Viruses enhanced the immunological outcome.
- Immunized monkeys challenged with SHIV showed a more rapid reduction of plasma viremia.

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

References Akridge et al. 1999

This study suggests that HIV-1-resistance in exposed and uninfected individuals is not only associated with the 32-bp deletion in the HIV-1 co-receptor CCR5, but can be related to HIV-1 specific CTL immunity.

**HXB2 Location** Env

Author Location gp120 (BRU)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Aladdin *et al.* 1999

• In vitro measurements of CTL-activity by Cr release assay in bulk culture showed no correlation between CTL-activity (gp120, Gag, Pol and Nef) and disease progression as measured by viral load, CD4 and time to death.

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Aladdin et al. 2000

 The administration of IL-2 caused an initial enhancement of CD4 cell counts that was accompanied by a decrease in CTL activity – IL-2 therapy did not reduce initial HIV viral load and viral replication was ultimately enhanced.

HXB2 Location Env

**Author Location** Env

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Jin et al. 1998a

- CTL precursor frequencies were determined in HIV-1 infected pregnant women, and higher CTLp frequencies to Pol and SF2 Nef, but not IIIB Nef, were found in non-transmitting mothers than in transmitting mothers – Nef CTL responses have been found in uninfected infants born to HIV+ women (Lazuriaga95);
- Very different CTLp frequencies were observed in env depending on whether IIIB, MN, RF, BK, or SF2 was used as antigen no association between env specific CTL and transmission was observed.

**HXB2 Location** Env

Author Location Env

**Epitope** 

Immunogen vaccine

Vector/Type: vaccinia HIV component: Env

Species (MHC)

Keywords review

References Zavala et al. 2001

- This paper is a review of vaccinia in the context of vaccines strategies that use different vectors to prime and boost, and emphasizes a unique capacity of vaccinia to very efficiently boost memory T-cell responses.
- HIV is discussed in the context of Gonazalo *et al.* 1999, where a V3 CTL epitope expressed in reFlu was boosted most effectively by vaccinia expressing the full Env.

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: ZF1 HIV component: complete genome

Species (MHC) macaque

References Akahata et al. 2000

- Rhesus macaques were vaccinated by i.m. injection with naked plasmid DNA carrying an HIV-1 complete genome vaccine, strain ZF1, with a mutated zinc finger in the nucleocapsid to prevent packaging.
- Env and Gag specific CTL but no antibody responses were induced in 2/4 vaccinated monkeys (MM145 and MM153)
- 2/4 monkeys (MM146 and MM143) produced antibodies against p24 and/or gp160, but no CTL response was detected.

- PBMC from all vaccinated monkeys produced IFN-gamma, 26/42 subjects who received CP vac-env-pro vaccine had a in response to HIV-1 gp160, indicating a Th response - this response was 5 times higher in MM145, the animal with the strongest CTL response.
- 4 weeks post-challenge with SHIV NM-3rN plasma viral loads of both MM145 and MM153 (with a homologous Env) decreased to near or below the detection limit.
- 6-8 weeks post-challenge with SHIV NM-3rN plasma viral loads of both MM146 and MM143 decreased near or below the detection limit.

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Young et al. 2001

- Addition of recombinant rec human IL12 (rhIL12) to cultures increased HIV-specific lysis of HIV-Gag, Pol and gp120 vaccinia expressed antigens (11/15 tested increased lysis by > 5%) if the culture was derived from HIV+ individuals who had CD4 cells/ul > 500.
- 2/10 individuals with <200 CD4 cells/ul, and 3/10 individuals with 200-500 CD4cells/ul, had an increase of >5% upon treatment of the culture with rhIL12, so a few individuals in late stage disease had CD8 cells that maintained responsiveness to rhIL12.

**HXB2 Location** Env

Author Location Env (subtype A, B, D)

**Epitope** 

Subtype A. B. D

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons

References Cao et al. 2000

- HIV-1 subtypes A and D dominate the Ugandan epidemic, and a vaccine trial using B clade antigen is underway – this study addresses relative levels of cross-reactive CTL responses in HIV infected Ugandans to A, D, and B clade recombinant vaccinia viruses expressing Gag, Env, Pol, RT or Nef from HIV-1 clades A, B, and D.
- Proteins corresponding to the subtype of the infecting strains tended to trigger higher levels of CTL response measured by percent specific lysis, but there was extensive inter-subtype cross-reactivity with B clade proteins and the co-circulating subtype.

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Immunogen vaccine

Vector/Type: canarypox, protein Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Env, Gag, Protease Adjuvant: MF59

Species (MHC) human

References AVEG022PT 2001

• Different HIV strains were used for different regions: MN (gp120), LAI (gp120, protease and gag), and SF2 gp120

- CTL response measured by Cr-release, while only 3/17 who were vaccinated with rec gp120 had a CTL response.
- A combination of a CP vac-env-pro vaccine with rec gp120 gave CD8+ T-cells in 62% of subjects, and NAbs in 91% of subjects.

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References White et al. 2001

· HIV-specific CTL activity was detected in the female reproductive tract of only 1/3 HIV-infected women who underwent a hysterectomy, although CTL could be identified in the PBMC of all three women.

**HXB2 Location** Env

**Author Location** Env (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression

References Jin et al. 2000a

- The CTL precursor level (CTLp) was measured in long term non-progressors (LTNP) with low viral load using limiting dilution analysis and measuring CTL against Env Gag and Pol expressed in vaccinia in autologous targets.
- · LTNPs have high memory CTL numbers and low viral load.

**HXB2 Location** Env

Author Location Env (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, rate of progression

References Jin et al. 2000a

- The CTL effector levels (CTLe) were compared in long term non-progressors (LTNP) with low viral load and in patients whose virus was well-suppressed by therapy, using a tetramer
- LTNPs have high memory CTLe numbers and low viral load, while HAART patients had low CTLe numbers and low viral load.

HXB2 Location Env

**Author Location** Env

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

Keywords review, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 2001

• This is a review that summarizes observations about HIVspecific CTL found in the HIV-1 exposed persistently seronegative (HEPS) population.

- The CTL responses assayed by ELISPOT and by CTL precursor frequencies by limiting dilution analysis indicate that CTL in HEPS individuals tend to be of a lower magnitude than in chronic HIV-1 infections the responses in HEPS cases are below the level of detection by tetramer assays.
- CD8+ CTL responses tend to be detectable in HEPS subjects only if they are recently exposed, and the response diminishes if exposure is reduced – it is not clear if there is a stable memory population in HEPS cases.
- CD8+ CTL responses in the HEPS population are associated with HIV-1 specific CD4+ T cell responses, assayed by proliferation assays, IL-2 secretion, and ELISPOT, and the authors consider the possibility that HIV-1-specific T-help responses improve the "quality" of the CD8+ response in HEPS individuals relative to HIV-1 infected individuals, who tend to have a poor HIV-1-specific T-help response.
- HIV-1 specific CD8+ CTL responses in HIV-1 infected individuals show reduced levels of perforin, and the T cells may not mature properly, and although similar studies have not been conducted in HEPS individuals this is considered as a possible difference in the CTL immune response in HEPS and HIV-1 infected people.

HXB2 Location Env Author Location Epitope Subtype B

Immunogen vaccine

Vector/Type: DNA HIV component: Env, Gag, Pol

Species (MHC) mouse
Keywords review

**References** Nabel 2002

- Env DNA constructs were designed that were codon optimized for human genes, express Env in the absence of the regulatory protein Rev, both increasing Env expression levels, deletions in the cleavage site and in the fusion domain. These constructs increased Ab responses to Env, while not diminishing CTL responses, when injected into mice.
- Removing N-linked glycosylation sites did not alter the humoral or cellular immune responses to this HIV protein, as has been seen in analogous SIV experiments.

**HXB2 Location** Env

**Author Location** 

Epitope Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS), responses in children, mother-to-infant transmission

References De Maria et al. 1994; Kuhn et al. 2002

- 6/24 HIV uninfected infants (ages 15-50 months) born to HIV+ mothers had HIV-1 specific CTL responses to vacciniaexpressed Nef, Gag/Pol, Env.
- Reviewed in Kuhn et al. [2002].

**HXB2 Location** Env **Author Location** 

Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, HIV exposed persistently seronegative (HEPS), responses in children, mother-to-infant transmission, rate of progression

References Kuhn et al. 2002; Wasik et al. 1999

- In HIV-infected infants HIV-specific, CTL responses were not detectable in cord blood or in PBMC collected shortly after birth and were absent or remained very low in 3 infants with a rapidly progressive disease. For those who progressed more slowly, the HIV-specific CTL activity varied.
- The two infants with high levels of Env peptide-stimulated IL-2 responses had the highest CTLp frequencies.
- Stronger responses were detected after initiation of the antiretroviral therapy.
- Two babies that were not infected though born to HIV+ mothers had detectable though low HIV-specific CTLp responses to Env (1/2), Pol (2/2), Gag (1/2) cord blood and transiently in PBMC after birth.
- Reviewed in Kuhn et al. [2002].

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS), responses in children, mother-to-

infant transmission

References Aldhous et al. 1994; Kuhn et al. 2002

- Six of nine HIV vertically infected infants had HIV-1 specific CTL responses to vaccinia expressed Tat (4/6), Pol (6/6), Env (1/6), or Gag (1/6), but not all responses were detected at all time points.
- Two of eleven babies that were not infected though born to HIV+ mothers had detectable responses Tat (1/2), Pol (2/2), Gag (1/2).
- Reviewed in Kuhn et al. [2002].

HXB2 Location Env

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS), responses in children, mother-to-infant transmission

References Kuhn et al. 2002; McFarland et al. 1994

- Only 9% of HIV+ infants had HIV-specific CTL against Env or Gag in unstimulated PBMC. After CD3 stimulation of PBMC, Gag and Env specific CTL were found in PBMC from 91% and 78% of HIV-infected children, respectively, with high precursor frequencies.
- 2/9 babies that were not infected though born to HIV+ mothers had detectable responses to Env.

• Reviewed in Kuhn et al. [2002].

**HXB2 Location** Env **Author Location Epitope** Subtype B Immunogen HIV-1 infection Species (MHC) human

Keywords epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.

**HXB2 Location** Env **Author Location Epitope** Immunogen HIV-1 infection Species (MHC) human Keywords HAART, ART

References Trabattoni et al. 2002

• CD8+ T-cells that were stimulated by HIV-1 Env expressing targets from 25 HIV+ patients receiving ART and 17 ART-naive patients were compared. CTL from the individuals receiving ART showed increased TNFalpha production and a reduction of perforin and granzyme expressing CTL, suggesting a functional defect in ART-treated individuals, and a potential benefit of immunomodulants during therapy.

**HXB2 Location** Env **Author Location (HXB2)** 

**Epitope** Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression References Edwards et al. 2002

- 96% (26/27) chronically infected HIV-1 infected patients elicited gamma-IFN CD8+ T-cell responses against Gag.
- Nef and/or Pol CTL responses were detected in 86% of the
- The magnitude and breadth of Gag and p24 T-cell responses correlated with absolute CD4 counts, and inversely correlated with viral load.
- Pol and Int CTL responses correlated positively with absolute CD4+ T-cell count.

• Nef and Env responses did not correlate with either CD4 counts or viral load.

**HXB2 Location** Env **Author Location** Env **Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB HIV component: gp160, Rev Adjuvant: cationic liposome, GM-CSF, IL-2

Species (MHC) mouse

Keywords Th2

References Ishii et al. 2001

- Vaccination route of HIV-1 DNA immunization with gp160 and Rev genes was compared including intranasal (i.n.), intramuscular (i.m.), and topical application of DNA directly on the skin after elimination of keratinocyte layers using a strong adhesive. Topical exposure resulted in high level CTL responses, IFNgamma and IL-4 production, and delayed type hypersensitivity (DTH). Topical application favored Th2 responses.
- DNA delivered topically with adjuvant-like cationic liposomes gave a stronger response than DNA alone, and coadministration of the DNA vaccine with IL-12 and GM-CSF expression vectors enhanced cytotoxic activity and DTH.

**HXB2 Location** Env **Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, dendritic cells

References Larsson et al. 2002b

• Autologous mature dendritic cells with rec vaccinia expressing Gag, Pol, Nef and Env could amplify CD8+ T-cell Elispot responses 4-38 fold in five HIV+ patients on successful HAART treatment, relative to autologous monocytes. Some weak responses could only be detected using mature dendritic cells as APCs, and this approach could be useful for detection of low frequency memory cells.

**HXB2 Location** Env

**Author Location (IIIB)** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords immunotherapy References Trickett et al. 2002

• Conditions were optimized for ex-vivo expansion of CD8+ and CD4+ T-cells with the goal of functional T-cell production for autologous immunotherapy. 10,000-fold expansions were obtained in 14 days with optimized concentrations of IL-2, anti-CD3 and anti-CD28 coated microspheres, and decreasing amounts of serum over the first 8 days.

**HXB2 Location** Env

**Author Location (IIIB)** 

**Epitope** 

Subtype B

Immunogen HIV-1 and HCV co-infection

Species (MHC) human

Keywords rate of progression

References Lauer et al. 2002

- HIV-1 and HCV immune responses were studied in 22 individuals who were co-infected with HIV-1 and hepatitis C virus (HCV). IFNgamma production was measured in an Elispot assay of CD8+ T-cells using targets expressing either Gag, RT, Env and Nef in a vaccinia construct, or one of seven HCV proteins.
- All 22 patients targeted at least one protein. 20/22 patients recognized RT, 17/22 patients recognized Gag, 13/22 subjects recognized Env and 11/22 patients recognized Nef. Robust CTL activity was independent of disease progression or viral load
- Despite high HCV viral loads, very few HCV CD8+ T-cell Elispot responses were detected. In a control HCV infected person who did not have HIV-1, strong anti-HCV responses were mounted.
- HIV-specific CD4 proliferative responses were detected in 9/17 coinfected patients, but no HCV responses were detected.

HXB2 Location Env

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords responses in children

References Luzuriaga et al. 1995

- 2/3 infants infected in utero had detectable HIV-1 Gag and Env specific CTL responses, one by 4 months, one by 11 months of age. Levels of the responses varied at different time point. Pol responses were not detected.
- 2/4 infants infected intrapartum had detectable responses, one note until 11 months, one not until 42 months.
- HIV-specific CTL were not detected in ten HIV- infants that were born to HIV+ mothers.

**HXB2 Location** Env

**Author Location** 

Epitope

Immunogen vaccine

*Vector/Type:* canarypox prime with gp120 boost *HIV component:* Env, Gag

Species (MHC) human

References Gupta et al. 2002

• A safety and immunogeniticity study of a vaccine dosing schedule was studied in a trial conducted in high and low risk study subjects. There was a 76% cumulative probability of detecting a Gag or Env CTL response by day 728.

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, responses in children

References Scott et al. 2001

• CTL responses before and after initiation of ART were studied in 13 HIV-1 vertically infected infants <6 months of age, and 4 that were >6 months of age.

- Before ART 2/13 infants <6 months of age showed IFNgamma Elispot CD8+ T-cell responses, one to Nef and one to Env and Nef, and these responses became undetectable after successful therapy—3 infants were coinfected with CMV and all 3 had CMV-specific CD8+ T-cell responses.
- One older infant, at 23 months, had CTL responses against all for proteins tested, Gag, Pol, Nef and Env, and had the lowest plasma viremia of the study group. 3/4 infants older than 6 months of age responded to either Nef or Pol.
- Administration of ART over 48 weeks broadened the HIV-1specific CTL response in 2/4 of the older children that were incomplete responders.

**HXB2 Location** Env

**Author Location** (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Ortiz et al. 2001

- Immune responses in eight chronically HIV-1 infected patients undergoing HAART therapy structured treatment interruptions (STI) were studied. STI boosted HIV-1 specific CTL responses and elevated CTL responses were maintained up to 22 weeks after the last treatment interruption, but viral load rebound to pretreatment levels and CD4 T-cell count decline was observed. CD8 responses in PBMC were measured by cytokine flow cytometry with gp160, Gag p55, RT-Pol and Nef expressed in vaccinia.
- One of seven subjects with a detectable NAb response had an augmented neutralization titer in response to STI.

**HXB2 Location** Env

**Author Location (SF2)** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC B\*3501, A\*2402, B\*5101, A\*3303

Keywords class I down-regulation by Nef

References Tomiyama et al. 2002

 Nef down-regulates class I molecules, and the killing activity of HLA B\*3501, A\*2402, B\*5101 and B\*3303-restricted HIV-1epitope specific CTL clones was inhibited by an HIV-1 strain carrying Nef, relative to a Nef-deleted virus; while Nef-induced HLA class I down-regulation inhibited lysis, it did not abolish cytokine production by HIV-1-specific CD8+ T-cells.

HXB2 Location Env

Author Location Env (gp160) (IIIB)

**Epitope** 

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade NL43

HIV component: Env

Species (MHC) macaque

References Akahata et al. 2003

• Four monkeys were injected i.m. with a SHIV plasmid (SHIV-NM-3rn ZF1\*) which encodes all viral proteins driven by the SIV LTR promoter. Infectivity is prevented by the introduction of mutations within the zinc-finger motifs of the nucleocapsid (NC) that prevents RNA packaging. An original NC ZF1 mutant plasmid was constructed using NL43 (Akahata 275:116-124 (2000) – the SHIV construct was made as an alternative to get improved expression in macaques using an SIV promotor. CTL were detected by lysis of HIV-1 Env IIIB or SIV Gag mac239 expressing expressing target cells, and a T cell proliferative response to Env was observed. Env-directed antibodies were detected by ELISA. All vaccinated macaques had a low peak viral loads that fell below the level of detection within 6 weeks post-challenge with autologous SHIV SHIV-NM-3rn.

**HXB2 Location** Env

**Author Location** Env (MN)

**Epitope** 

Subtype B

Immunogen SIV infection, SHIV infection

Species (MHC) macaque

**Assay type** CD8 T-cell Elispot - IFNγ

Keywords assay standardization/improvement

References Calarota et al. 2003

- The sensitivity of gamma INF Elispot assays can be enhanced for the detection of low frequency responses, like after ART, by adding IL-15 to the assay.
- CD8+ T-cells from SHIV and SIV infected macaques with peptide pools from Gag and Env were used to test this system.

**HXB2 Location** Env

Author Location Env

**Epitope** 

Subtype multiple

**Immunogen** 

Species (MHC) human

**Assay type** Flow cytometric T-cell cytokine assay

**Keywords** subtype comparisons References Currier et al. 2003

- CD8-cellular immune responses from 21 HIV-1 infected patients from Kenya infected with subtype A, C, D, and unique recombinants were studied for cross-recognition of Gag, Env, and Nef vaccinia-expressed proteins representing subtypes A-H, including CRF01.
- Both subtype-specific and cross-reactive CTL were observed, with a skewing of responses towards the infecting subtype in the nine subjects with full length sequence available. The magnitude of the responses to Gag were the highest, less to Nef, and still less to Env.
- For Gag, 8/21 subjects responded to at least 7/8 different subtype proteins, 7 had a mixed response, recognizing some subtypes but not others, and 6 responded to only one or none of the different subtype proteins. For Env, 4/19 subjects responded to at least 7/8 subtypes, 8 had a mixed response, and 7 responded to one or none. For Nef, 5/19 subjects responded to at least 7/8 subtypes, 8 had a mixed response, and 6 responded to one or none.

**HXB2 Location** Env

Author Location Env (HIV-1 IIIB)

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

Assay type Cytokine production

Keywords HIV exposed persistently seronegative (HEPS)

References Fowke et al. 2000

- A cohort of Nairobi sex-workers were defined as resistant to HIV-infection by virtue of remaining seronegative despite repeated high risk exposures. 24 were tested for HIV specific T-helper responses determined by IL-2 production in vitro in response to gp120 peptides or soluble gp120 protein.
- 7/17 resistant women showed IL-2 stimulation which was greater than or equal to 2.0, and specific CTL responses were detected in 15/22 resistant women as compared to 0/12 of the control low-risk subjects.

**HXB2 Location** Env

Author Location gp160 (MN)

**Epitope** 

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade MN HIV component: gp160 Adjuvant: IL-12

Species (MHC) mouse

Assay type proliferation, CD8 T-cell Elispot - IFNγ,

Chromium-release assay

Keywords adjuvant comparison

References Chattergoon et al. 2004

• pIL-12 used as adjuvant significantly increases the number of Ag-specific CD8+ T-cells and a sustained memory response. Also, the splenocytes from mice that received pIL-12 were shown to proliferate to a much higher extent. Mice immunized with a plasmid expressing the influenza A/PR8/34 HA gene and pIL-12 were better able to control the infection.

HXB2 Location Env

Author Location Env

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ, Intracellular cytokine staining, Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, immune dysfunction

References Trabattoni et al. 2004

Reduced perforin-and granzyme- containing Env-specific CD8+ T cells were observed in ART treated individuals indicating that antiretroviral drugs might directly interfere with the production of perforin and granzymes, inhibiting CTL killing. Immunomodulants may be needed to enable CTL to become fully functional during ART.

**HXB2 Location** Env

**Author Location** gp140

**Epitope** 

Immunogen vaccine

Vector/Type: protein, peptide in liposome Strain: B clade IIIB HIV component: gp140, gp160, oligomeric gp140 Adjuvant: CpG immunostimulatory sequence (ISS), liposome

**Species (MHC)** mouse **Donor MHC** H-2d

Assay type Cytokine production, proliferation, Chromium-release assay

**Keywords** Th1, Th2, adjuvant comparison, vaccine antigen design

References Rao et al. 2004

• Administration of ogp140 in liposomes containing lipid A (LA) induces high antibody titers which are increased by adding CpG ODN. Priming and boosting of BALB/c mice with ogp140+LA induces mixed Th1/Th2 immune response, while adding CpG ODN switches the immune response to a Th1 type. Mixing ogp140 with lipomoses containing lipid A yielded excellent proliferative and CTL specific responses; CpG did not affect CTL responses. The antigen did not need to be encapsulated in the liposome to induce strong responses with LA as an adjuvant.

HXB2 Location Env Author Location Epitope

Subtype CRF02\_AG

Immunogen vaccine

Vector/Type: virus-like particle (VLP), DNA prime with modified vaccinia Ankara (MVA) boost Strain: CRF02 IC0928 HIV component: Env, Gag, Pol

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining Keywords vaccine-specific epitope characteristics, vaccine antigen design

References Ellenberger et al. 2005

 Macaques were given a Gag-Pol-Env DNA prime followed by an MVA boost. Two DNA constructs were compared, one that resulted in mature VLPs with processed Gag (IC48) and one that had a point mutation in Gag that resulted in immature VLPs (IC1-90). IC48 DNA vaccinations, which produced mature VLPs, yielded 2-fold stronger T-cell responses with greater breadth. CD4 T-cells responded to 3-fold more peptide pools than did CD8.

**HXB2 Location** Env **Author Location** 

**Epitope** 

**Subtype** CRF02\_AG **Immunogen** vaccine

Vector/Type: virus-like particle (VLP), DNA prime with modified vaccinia Ankara (MVA) boost Strain: CRF02 IC0928 HIV component: Env, Gag, Pol

Species (MHC) macaque

**Assay type** T-cell Elispot, Intracellular cytokine staining **Keywords** vaccine-specific epitope characteristics, vac-

cine antigen design

References Ellenberger et al. 2005

 Macaques were given a Gag-Pol-Env DNA prime followed by an MVA boost. Two DNA constructs were compared, one that resulted in mature VLPs with processed Gag (IC48) and one that had a point mutation in Gag that resulted in immature VLPs (IC1-90). IC48 DNA vaccinations, which produced mature VLPs, yielded 2-fold stronger T-cell responses with greater breadth. CD4 T-cells responded to 3-fold more peptide pools than did CD8.

HXB2 Location Env Author Location

Epitope Subtype B

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: Other HIV component: Env, Gag, Pol, Rev, Tat, Vif,

Vpr

Species (MHC) macaque

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining

**Keywords** vaccine-specific epitope characteristics, vaccine antigen design, characterizing CD8+ T

References Sadagopal et al. 2005

- 22/23 macaques that were immunized with a DNA prime SHIV-89.6 and boosted with rMVA showed successful control of viremia, with low or undetectable viral loads and normal CD4 counts 200 weeks postchallenge. IFN-gamma producing T cells were found in unexpectedly low breadths and frequencies. T-cell responses were stable over time and maintained their production of IFN-gamma and IL-2. Long-term control was found in macaques of diverse histocompability types. The CD8 T cells seemed to have the most impact on well-contained chronic infections in the vaccinated and challenged animals.
- Both CD4 and CD8 responses were found to the SIV Gag and HIV Env proteins, 60% of CD8+ epitopes were in p27, and 80% of CD4+ epitopes.

## II-B-23 Nef CTL/CD8 + epitopes

HXB2 Location Nef (1–16)

**Author Location** Nef (1–16)

Epitope MGGKWSKSSIVGWPAV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Nef (5–13)

Author Location Nef (5–13 HXB2) Epitope WSKSSIIGW

Epitope name WW9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2501)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords escape, immune evasion, optimal epitope,

HIV-1

References Liu et al. 2006

- T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
- This is a newly defined epitope. Positions 4 and 6 in the epitope had potentially experienced positive selection. WSKtSIIGW and WSKSSmIGW escape variants were found.

HXB2 Location Nef (11-20)

**Author Location Nef** 

Epitope VEWPAVRERM

Epitope name VM10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 2, VgWPAVRERM, was found not to correspond to the most polymorphic residue in the epitope. This is a novel unmapped epitope.

**HXB2 Location** Nef (13–20)

**Author Location** Nef (13–20 LAI)

Epitope WPTVRERM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Keywords optimal epitope

References Frahm et al. 2007; Goulder et al. 1997g

• C. Brander notes this is a B\*0801 epitope.

**HXB2 Location** Nef (13–20) **Author Location** Nef (HXB2)

Epitope WPTVRERM

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** (B\*0801)

Keywords class I down-regulation by Nef

References Peng & Robert-Guroff 2001

B\*1801, B\*5101,
 Deletion of the 19 N-terminal amino acids from Nef including the myristolation signal eliminates Nef-induced downregulation of MHC class I and CD4 molecules. Such a construct has the potential to serve as a more potent immunogen. The known T-cell epitopes that that would be disputed by this deletion are minimal, including the HLA-B8 CTL epitope WPTVRERM.

HXB2 Location Nef (13-20)

Author Location (C consensus)

Epitope WPAIRERM

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location Nef (13-20)

**Author Location** Nef

Epitope WPTVRERM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

**Donor MHC** A\*0101, B\*0801

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , HLA binding

**Keywords** escape, acute/early infection

References Milicic et al. 2005

• CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient. In an early acute infection of the donor, the precise timing of transmission determines the viral variants transmitted.

The recipient mounted an acute CTL response to this epitope, and the escape variant wpAvrKrm emerged in the recipient soon after.
 The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preserve on after.

HXB2 Location Nef (13–20)
Author Location (C consensus)
Epitope WPAIRERM
Subtype C
Immunogen HIV-1 infection
Species (MHC) human (B\*0801)
Country South Africa.

**Assay type** CD8 T-cell Elispot - IFNγ **Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- WPAIRERM is an optimal epitope.

HXB2 Location Nef (13–20) Author Location Nef (13–20 LAI) Epitope WPTVRERM

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8)

References Goulder et al. 1997g

Unusual epitope for HLA-B8, but compatible with crystal structure predictions.

HXB2 Location Nef (13–20)
Author Location Nef (13–20)
Epitope WPTVRERM
Immunogen HIV-1 infection
Species (MHC) human (B8)
Keywords immunodominance
References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the HLA A2+ was HLA A\*0201, A31, B8, B51 and responded to this epitope as well as seven others.

HXB2 Location Nef (13–20)
Author Location Nef (13–20 SF2)
Epitope WPTVRERM
Immunogen HIV-1 infection
Species (MHC) human (B8)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

 Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.

- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B8+ individuals that had a CTL response to this epitope broken down by group: 1/3 group 1, 0/3 group 2, and 1/2 group 3.

HXB2 Location Nef (13–20) Author Location Nef (13–20) Epitope WPTVRERM Immunogen HIV-1 infection Species (MHC) human (B8)

References Day et al. 2001

• B8-restricted CTL accounted for about 1/3 of the total CTL response in one individual.

HXB2 Location Nef (13–20)
Author Location Nef (13–20)
Epitope WPTVRERM
Epitope name WM8
Immunogen HIV-1 infection
Species (MHC) human (B8)

**Donor MHC** A\*03, A\*31, B\*08, B\*15, Cw\*04, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection, reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs.
   Escape variants were commonly detected in maternal plasma.
   Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- The variant WnTVRERM was present in 10/10 clones from a B8+ mother, was transmitted to her B8- infant, and present in 10/10 clones at months 2, 4, and 15.

HXB2 Location Nef (19–27)
Author Location Nef (19–27)
Epitope RMRRAEPAA
Immunogen HIV-1 infection
Species (MHC) human (B62)
Keywords optimal epitope
References Frahm et al. 2007

HXB2 Location Nef (19–27) Author Location Nef (19–27) Epitope RMRRAEPAA Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B62)

Donor MHC 1261: A\*0201, A29, B58, B62, Cw\*0304,

Cw\*1601

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

HXB2 Location Nef (22-30)

Author Location Nef (22–30)

Epitope RAEPAADGV

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*6901)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, computational epitope

prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

HXB2 Location Nef (29-37)

Author Location Nef (29–37)

Epitope GVGAASRDL

Subtype C

**Immunogen** HIV-1 infection **Species (MHC)** human (A\*6901)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

HXB2 Location Nef (37–45)

**Author Location** Nef (37–45)

Epitope LEKHGAITS

Immunogen HIV-1 infection

**Species (MHC)** human (B\*4001) **Keywords** optimal epitope

References Frahm et al. 2007

HXB2 Location Nef (37–45)

**Author Location** Nef (37–45)

Epitope LEKHGAITS

Immunogen

Species (MHC) human (B50)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Nef (42–50)

**Author Location** Nef (44–52 HXB3)

Epitope ALTSSNTAA

Immunogen vaccine

Vector/Type: DNA, peptide Strain: B clade HXB3 HIV component: Nef Adjuvant:

Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (A\*0201)

**Keywords** binding affinity, computational epitope predic-

References Sandberg et al. 2000

- Ten Nef 9-mer peptides were predicted to have a strong binding affinity with HLA-A\*0201 of these, four did bind strongly by a T2 class I stabilization assay, several others bound weakly.
- A CTL immune response to only 3/10 peptides was detected by a 51Cr-release assay after immunization of HLA-A201 transgenic mice with either nef DNA under the control of a CMV promotor, coated on gold particles delivered to abdominal skin by gene gun.
- ALTSSNTAA was also tested by subcutaneous injection of Nef peptides in Freund's adjuvant.
- ALTSSNTAA bound weakly to HLA-A2, but it had the strongest CTL response among the three elicited by the DNA vaccine and a strong response to the peptide vaccination.

HXB2 Location Nef (42–50)

**Author Location** Nef (42–50)

**Epitope ALTSSNTAA** 

**Epitope name** Nef42-50

Immunogen HIV-1 infection

Species (MHC) human, humanized mouse (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords responses in children, immunodominance,

characterizing CD8+ T cells

References Chandwani et al. 2004

 Responses to five HLA-A2 presented epitopes in Nef were characterized in a population of 19 HIV infected adults and 21 children. The CD8 T-cell response to Nef was stronger and broader in adults than children, averaging 652 and 87 SFCs/10<sup>6</sup> PBMC, and 2.6 and 0.9 recognized epitopes per person, respectively.

• This is one of three novel Nef epitopes previously identified in HLA-A2 transgenic mice, shown to induce CD8 T-cell response in humans. It was not the immunodominant response.

HXB2 Location Nef (48–56)

**Author Location** Nef (58–66 JRFL)

**Epitope** TAATNADCA

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade JRFL

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Liang et al. 2002

- BALB/c, C3H/HeN and C57BL/6 mice were given intramuscular immunization with Nef DNA constructs – C57BL/6 responded to this epitope.
- The Nef mutant that lacked the myristylation site (G->A) at position 2, and the dileucine motif (L -> A at positions 174 and 175) was impaired in terms of its ability to elicit induction of Nef-specific CD4+ and CD8+ T-cell responses. The myristylation site is critical for Nef membrane localization and function, and the di-leucine motif for the down-regulation of surface CD4 molecules, and the mutation of these regions could yield a safer vaccine.
- N-terminal addition of human tissue plasminogen activator (TPA) to Nef, enhanced CD8+ T-cell responses and could compensate for the G2A, L174A, L175A mutations – this enhanced immunogenicity correlated with enhanced levels of protein expression in transfected cells.

HXB2 Location Nef (50–58)

**Author Location** Nef (50–)

Epitope ATNADCAWL

**Epitope name** Nef50

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Nef Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (MHC) human (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords binding affinity, subtype comparisons, com-

putational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was a low A2-binder that did not induce CTL or CD8+ T-cell IFN gamma responses in mice. Responses were detected in 2/17 HIV+ HLA-A2 subjects.

HXB2 Location Nef (62–81)

Author Location Nef (61-80)

Epitope EEEEVGFPVTPQVPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

 HIV-specific CTL lines developed by ex vivo stimulation with peptide.

HXB2 Location Nef (62–81)

Author Location Nef (61-80 SF2)

Epitope EEEEVGFPVTPQVPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Twelve subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- Two of these 12 had CTL response to this peptide.
- The responding subjects were HLA-A11, A24, B8, B35, and HLA not determined.

HXB2 Location Nef (62–81)

Author Location Nef (61-80 SF2)

Epitope EEEEVGFVTPQVPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

 CTL expanded ex vivo were later infused into HIV-1 infected patients.

HXB2 Location Nef (62–81)

Author Location Nef (SF2)

Epitope EEEEVGFPVTPQVPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2001a

- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- Nef peptides PQVPLRRMTYKAAVDLSHFL, KAAVDLSH-FLKEKGGLEGLI and EEEEVGFPVTPQVPLRPMTY were recognized and the first two share KAAVDLSHFL (a Cw8 epitope), the first and last share PQVPLRPMTY.

HXB2 Location Nef (64–74)

**Author Location** Nef (C consensus)

Epitope GEVGFPVRPQV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B45)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords escape, reversion, viral fitness

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- People who carried B45 tended to carry a variant of this epitope, while people who did not almost always carried the consensus form.
- B\*4501 was one of the HLA types associated with having a high viral load.

HXB2 Location Nef (66-80)

Author Location Nef (66-80 BRU)

 ${\bf Epitope} \ \ {\tt VGFPVTPQVPLRMT}$ 

Immunogen HIV-1 infection

Species (MHC) human (A1, B8)

References Hadida et al. 1992

• HIV-1 specific CTLs detected in lymphoid organs of HIV-1 infected patients.

HXB2 Location Nef (66–80)

**Author Location** Nef (64–78)

Epitope VGFPVTPQVPLRMT

Immunogen HIV-1 infection

Species (MHC) human (A1, B8)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location Nef (66–97)

Author Location Nef (66–97 LAI)

Epitope VGFPVTPQVPLRPMTYKAAVDLSHFLKEKGGL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long amino acid peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees – 5/10 reacted to this Nef peptide.
- 9/12 tested mounted a CTL response to at least one of the six peptides; each of the six peptides elicited a CTL response in at least one individual.
- 5/12 tested had an IgG response to this peptide.

HXB2 Location Nef (67-81)

**Author Location** Nef (67–81)

**Epitope** GFPVRPQVPLRPMTY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Nef (68–76)

**Author Location** Nef (68–76)

Epitope FPVTPQVPL

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Keywords optimal epitope

References Frahm et al. 2007

HXB2 Location Nef (68–76)

**Author Location** Nef (103–111)

Epitope FPVRPQVPL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Assay type Other

Keywords HLA associated polymorphism

References Boutwell & Essex 2007

- All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.
- FPVRPQVPL was a previously defined B\*0702 presented epitope that encompassed a B\*07 associated polymorphism, FPVr-PQVPL, in the fourth position.

HXB2 Location Nef (68–76)

**Author Location** Nef (68–76)

Epitope FPVRPQVPL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*35)

**Donor MHC** A\*03, A\*24, B\*35, B\*40

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** acute/early infection, variant crossrecognition or cross-neutralization, superinfection

References Yang et al. 2005b

- An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- This epitope FPVRPQVPL was identical in the initial and superinfecting strains, and the CTL response persisted in the patient before and after superinfection.

HXB2 Location Nef (68–76)
Author Location Nef (72–80 SF2)
Epitope FPVRPQVPL
Immunogen HIV-1 infection
Species (MHC) human (B\*3501)
References Tomiyama et al. 1997

- A CTL clone responsive to this epitope was obtained.
- 3/7 B35-positive individuals had a CTL response to this epitope.
- An R to T substitution at position 4 abrogates specific lysis, but not binding to B\*3501.

HXB2 Location Nef (68–76)
Author Location Nef (72–80)
Epitope FPVRPQVPL
Immunogen HIV-1 infection
Species (MHC) human (B\*3501)
References Tomiyama et al. 2000a

- CD8+ T-cells that bound one of six HIV-specific B\*3501-epitope tetramers did not express CD28 or CD45A.
- A significant increase in CD28-CD45RA- cells and a decrease of CD28+CD45RA+ cells was observed in chronically HIV-1infected individuals relative to healthy individuals.
- CD28-CD45RA- cells are likely to be effector cells and have high levels of perforin in their cytoplasm.
- The mean percentage of total CD28- CD8+ cells in chronically infected HIV-1-infected patients was 76.6% in comparison to HIV-1-uninfected individuals (40.6%)

HXB2 Location Nef (68–76)
Author Location Nef (72–80 SF2)
Epitope FPVRPQVPL
Immunogen HIV-1 infection
Species (MHC) human (B35)
References Shiga et al. 1996

• Binds HLA-B\*3501.

HXB2 Location Nef (68–76)
Author Location (SF2)
Epitope FPVRPQVPL
Immunogen HIV-1 infection
Species (MHC) human (B35)

**Keywords** rate of progression **References** Kawana *et al.* 1999

- HLA B35 is associated with rapid disease progression.
- The sequences of 9 previously described HIV-1 B35 CTL epitopes were obtained in 10 HLA B35+ and 19 HLA B35- individuals.
- 3/9 CTL epitopes had substitutions that were more common in B35+ individuals than in B35- individuals, but this was one of the six that had no B35 associated pattern of mutation.

HXB2 Location Nef (68–76)
Author Location Nef (66–74)
Epitope FPVRPQVPL
Immunogen HIV-1 infection
Species (MHC) human (B35)
References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location Nef (68–76)
Author Location Nef (68–76 BRU)
Epitope FPVTPQVPL
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (B35)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- FPVTPQVPL was recognized in 1/13 (8%) of individuals with HLA B7, and 1/12 (8%) of individuals with HLA B35. It was a high affinity HLA binder.

HXB2 Location Nef (68–76) Author Location Nef (68–76) Epitope FPVTPQVPL

Immunogen in vitro stimulation or selection

Species (MHC) human (B7)

Keywords binding affinity, dendritic cells, Th1

References Wilson et al. 1999b

- Dendritic cells are the most potent for priming T cell responses
   DCs can stimulate autologous CTL responses from T cells cultured from HIV negative donors.
- Th1-biasing cytokines IL-12 or IFN alpha enhance CTL responses *in vitro* whether the epitope is delivered by pulsing from peptide, or expressed from within.
- B7 and A2 Nef epitopes were studied FPVTPQVPL has a high affinity for B7.

**HXB2 Location** Nef (68–76) **Author Location** Nef (68–76)

Epitope FPVTPQVPL Immunogen HIV-1 infection Species (MHC) human (B7)

**Keywords** rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

HXB2 Location Nef (68–76)

Author Location Nef (68–76 BRU) Epitope FPVTPQVPL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- FPVTPQVPL was recognized in 1/13 (8%) of individuals with HLA B7, and 1/12 of individuals with HLA B35. It was a high affinity HLA binder.

**HXB2 Location** Nef (68–76)

**Author Location** Nef (68–76)

Epitope FPVTPQVPL

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

 CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.

- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. Also, none of 4 individuals had detectable responses to this epitope after STI.

HXB2 Location Nef (68–76)

Author Location Nef (68-76)

Epitope FPVTPQVPL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (B7 supertype)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

Keywords vaccine-induced epitopes, variant cross-

recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+T-cell epitopes was induced after a fourth boost injection. This epitope was one of the most frequently recognized of the 31 epitopes that the vaccinated volunteers responded to.

HXB2 Location Nef (68–77)

**Author Location** Nef (68–77 LAI)

Epitope FPVTPQVPLR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

**Keywords** optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*0702 epitope.

HXB2 Location Nef (68–77)

**Author Location** Nef (68–77 LAI)

Epitope FPVTPQVPLR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Haas et al. 1996

 There was a high degree of variation in three CTL epitopes in Nef in four slow and non-progressors, and variant specific CTLs arose over time to eliminate variants, indicating immune selection.

HXB2 Location Nef (68–77)

**Author Location** Nef (subtype B)

Epitope FPVTPQVPLR

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

**Keywords** HIV exposed persistently seronegative (HEPS), escape

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- FPVTPQVPLR was recognized in 1 of the 6 women (ML1203), and the response was present in the last available sample prior to seroconversion, 7 months.
- 20/20 sequences of the infecting strain had no substitutions in this epitope, all were FPVTPQVPLR, so there was no evidence for escape.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized in 1/22 HEPS sex worker controls, ML851.

HXB2 Location Nef (68–77)
Author Location Nef (66–75)
Epitope FPVRPQVPLR
Immunogen HIV-1 infection
Species (MHC) human (B7)
References Ferrari et al. 2000

 One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location Nef (68–77)
Author Location Nef (68–77 SF2)
Epitope FPVTPQVPLR
Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B7+ individuals that had a CTL response to this epitope broken down by group: 0/4 group 1, 0/3 group 2, and 1/1 group 3.

HXB2 Location Nef (68–77) Author Location Nef (68–77) Epitope FPVTPQVPLR Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (B7)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused
  on different epitopes with HLA presenting molecules that have
  previously been associated with reduced risk of infection, and
  there was a shift in the response in the HEPS women upon late
  seroconversion to epitopes recognized by the HIV-1 infected
  women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Subject ML 1203 started with CTL responses to A\*6802 DTVLEDINL and to B7 FPVTPQVPLR prior to seroconversion, and upon seroconversion acquired additional responses to A\*6802 ETAYFILKL which became dominant, B7 TPGPG(V/I)RYPL, B7 IPRRIRQGL, and B7 SPRTLNAWV.

**HXB2 Location** Nef (68–77) **Author Location** Nef (68–77)

Author Location Net (00-77)

Epitope FPVTPQVPLR

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

**HXB2 Location** Nef (68–77)

**Author Location** Nef (68–76)

Epitope FPVTPQVPLR

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions

(STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or
   B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
   T-cell immune responses were examined in HIV-1 infected Ugandans. Similar levels of cross-clade responses for Gag, Env and Nef were observed. Higher frequencies of responses
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. Also, none of 4 individuals had detectable responses to this epitope after STI.

HXB2 Location Nef (68–77)

Author Location Nef (66–75)

Epitope FPVTPQVPLR

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 7 patients recognized this epitope.

**HXB2 Location** Nef (68–81)

**Author Location** Nef (82–95 HXB2)

Epitope FPVTPQVPLRMTY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Guimaráes *et al.* 2002

 Nef sequences were obtained from Brazilians to study epitope diversity in this geographic region—the HXB2 sequence is FPVTPQVPLRMTY, but fpvRpqvplrmty was observed in most Brazilian sequences regardless of the subtype (A, C, D and F).

**HXB2 Location** Nef (68–82)

**Author Location** Nef (73–82)

Epitope FPVRPOVPLRPMTYK

Subtype A, D

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*0201, A\*0301, B\*4501, B\*5802

Country Uganda.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, characterizing CD8+ T

cells

References Barugahare et al. 2005

- T-cell immune responses were examined in HIV-1 infected Ugandans. Similar levels of cross-clade responses for Gag, Env and Nef were observed. Higher frequencies of responses in conserved regions were found for Gag within a single clade, while areas with higher viral sequence variation had fewer detectable responses. Differential interclade sequence homologies between the Gag regions did not affect the respective level of cross-clade recognition.
- This sequence contains a known epitope (RPOVPLRPMTYK).
   The subject recognizing the peptide carries an HLA allele (A0301) of the known restriction, and the peptide is conserved in the autologous sequence.

HXB2 Location Nef (68–84)

**Author Location** Nef

Epitope FPVRPQVPLRPMTYKGA

Immunogen

Species (MHC) human

**Keywords** subtype comparisons **References** Jubier-Maurin *et al.* 1999

- 41 new HIV-1 strains describing envelope subtypes of HIV-1
   A-H were genetically characterized in the nef region 34 subtypes were classified in the same subtype in nef and env and 7 of the 41 strains were recombinants.
- This region was defined as a CTL epitope region that is conserved among HIV-1 M group subtypes.

HXB2 Location Nef (70-84)

**Author Location** Nef (70–84 HXB2)

Epitope VTPQVPLRPMTYKAA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 34% of the study subjects, and it was the second most frequently recognized peptide.

HXB2 Location Nef (71–79)
Author Location Nef (71–79 LAI)
Epitope TPQVPLRPM
Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*0702)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*0702 epitope.

HXB2 Location Nef (71–79) Author Location (C consensus) Epitope RPQVPLRPM

Subtype C

Immunogen HIV-1 infection
Species (MHC) human (B\*4201)
Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- RPQVPLRPM is an optimal epitope for both B\*4201 and B\*4202.

**HXB2 Location** Nef (71–79)

**Author Location** 

Epitope RPQVPLRPM

Epitope name RM9

Immunogen

Species (MHC) human (B\*4201)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B\*4201 epitope.

HXB2 Location Nef (71–79) Author Location (C consensus) Epitope RPQVPLRPM

Subtype C

Immunogen HIV-1 infection Species (MHC) human (B\*4202)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- RPQVPLRPM is an optimal epitope for both B\*4201 and B\*4202.

HXB2 Location Nef (71–79) Author Location Nef (71–79 BRU) Epitope TPQVPLRPM

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B35)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- TPQVPLRPM was recognized in 1/10 (10%) of individuals with HLA B7, and 1/10 (10%) of individuals with HLA B35. It was a moderate affinity HLA binder.

HXB2 Location Nef (71–79)

**Author Location** Nef (71–79 SF2)

Epitope TPQVPLRPM

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B7+ individuals that had a CTL response to this epitope broken down by group: 0/4 group 1, 0/3 group 2, and 1/1 group 3.

HXB2 Location Nef (71–79)

**Author Location** Nef (71–79)

Epitope TPQVPLRPM

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.

- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

**HXB2 Location** Nef (71–79)

**Author Location** Nef (71–79 BRU)

Epitope TPQVPLRPM

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- TPQVPLRPM was recognized in 1/10 (10%) of individuals with HLA B7, and 1/10 (10%) individuals with HLA B35. It was a moderate affinity HLA binder.

HXB2 Location Nef (71–79)

**Author Location** Nef (71–79)

Epitope TPQVPLRPM

Epitope name B7-TM9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7) Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 2/4 individuals had detectable responses to this epitope after STI.

HXB2 Location Nef (71-79)

**Author Location** Nef

**Epitope** TPQVPLRPM **Epitope name** B7-TM9(Nef)

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A32, B7, B14; A24, B7, B27

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals. Patient A displayed the greatest response to epitope B14-EL9(gp41), a strong response to B7-TL9(p24), and responses to B7-TM9(Nef) and A32-PW10(RT). Patient C displayed the greatest response to B27-KK10(p24), and in decreasing order also responded to A24-RW8(Nef), B7-IL9(gp41), A24-RL9(gp41), A24-YL8(gp41), and B7-TM9(Nef).

**HXB2 Location** Nef (71–79)

**Author Location** Nef (180–187)

Epitope TPQVPLRPM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

**Country** United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

HXB2 Location Nef (71–79)

**Author Location** Nef (71–79)

Epitope TPQVPLRPM

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: OS21

**Species (MHC)** human (B7 supertype)

Chromium-release assay

Keywords vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

• After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the most frequently recognized of the 31 epitopes that the vaccinated volunteers responded to.

HXB2 Location Nef (71–79) Author Location Nef (C consensus)

Epitope RPQVPLRPM

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B7, B81, B\*4201, B\*0702, B\*8101)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, cross-presentation by different HLA

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- People who carried either B07 or B81 tended to carry a variant of this epitope, while people who did not almost always carried the consensus form.
- B\*4201 may also present this epitope, as the allele is enriched in people who react with the peptide that contains the epitope, and it is known from the database to be also presented by B\*4201.

HXB2 Location Nef (71–81)

**Author Location** Nef (75–85 SF2)

Epitope RPQVPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

References Tomiyama et al. 1997

- A CTL clone responsive to this epitope was obtained.
- 4/7 B35-positive individuals had a strong CTL response to this epitope.
- An R to T substitution at position 1 abrogates specific lysis, but not binding to B\*3501.
- An R to H substitution at position 7 did not alter reactivity.

HXB2 Location Nef (71–81) **Author Location** Nef (75–85) Epitope RPQVPLRPMTY Immunogen HIV-1 infection Species (MHC) human (B\*3501)

References Tomiyama et al. 2000a

- CD8+ T-cells that bound one of six HIV-specific B\*3501epitope tetramers did not express CD28 or CD45A.
- Assay type proliferation, CD8 T-cell Elispot IFN $\gamma$ , A significant increase in CD28-CD45RA- cells and a decrease of CD28+CD45RA+ cells was observed in chronically HIV-1infected individuals relative to healthy individuals.
  - CD28-CD45RA- cells are likely to be effector cells and have high levels of perforin in their cytoplasm.
  - The mean percentage of total CD28- CD8+ cells in chronically infected HIV-1-infected patients was 76.6% in comparison to HIV-1-uninfected individuals (40.6%)

HXB2 Location Nef (71–81)

**Author Location** Nef (75–85 SF2)

Epitope RPQVPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Shiga et al. 1996

• Binds HLA-B\*3501.

**HXB2 Location** Nef (71–81)

**Author Location (SF2)** 

**Epitope** RPQVPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human (B35)

**Keywords** binding affinity, rate of progression, escape

References Kawana et al. 1999

- HLA B35 is associated with rapid disease progression.
- The sequences of 9 previously described HIV-1 B35 CTL epitopes were obtained in 10 HLA B35+ and 19 HLA B35- indi-
- 3/9 CTL epitopes had substitutions that were more common in B35+ individuals than in B35- individuals – only one of these reduced the binding of the peptide to B35 and was shown to be an escape mutation.
- rpgvplrpmtF was found in 9/10 of the B35+ individuals, none of the B35- individuals—the Y->F substituted peptide had a similar binding affinity with B35 and was recognized by a CTL clone equally with wildtype.

HXB2 Location Nef (71–81)

Author Location Nef (69-79)

Epitope RPQVPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location Nef (71–81)

**Author Location** Nef (71–81 BRU)

**Epitope** TPQVPLRPMTY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- TPQVPLRPMTY was recognized in 9/12 (75%) of individuals with HLA B7, and 5/10 (50%) of individuals with HLA B35. It was a moderate affinity HLA binder, and the C-term Y readily cleaved *in vitro*.

HXB2 Location Nef (71-81)

**Author Location** Nef

Epitope RPQVPLRPMTY

Subtype A, B, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (B51)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location Nef (71–81)

**Author Location** Nef (71–81 BRU)

Epitope TPQVPLRPMTY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35,

- and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- TPQVPLRPMTY was recognized in 9/12 (75%) of individuals with HLA B7, and 5/10 (50%) of individuals with HLA B35. It was a moderate affinity HLA binder, and the C-term Y readily cleaved *in vitro*.

HXB2 Location Nef (71–81)

**Author Location** Nef (71–81)

Epitope TPQVPLRPMTY

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: QS21

**Species (MHC)** human (B7 supertype)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes, variant cross-recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

- After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.
- A response was induced in one patient after immuniaation with lipopeptides alone (no adjuvant) after the third (W44) boost. A rPQVPLRPMTY variant was also recognized.

HXB2 Location Nef (72–81)

**Author Location** Nef (72–82)

Epitope PQVPLRPMTY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35, B51)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot granzyme B

**Keywords** Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- Three of nine patients responded to this peptide with GzB producing and IFN-gamma producing cells, and one additional with IFN-gamma producing cells.

HXB2 Location Nef (72–86)

**Author Location** Nef (72–86)

Epitope PQVPLRPMTYKGAFD

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Nef (72–91)

**Author Location** Nef (71–90 SF2)

Epitope PQVPLRMTYKAAVDLSHFL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1
- · Eleven subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- Three of these 11 had CTL response to this peptide.
- The responding subjects were HLA-A3, A32, B51, B62; HLA- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted A11, A24, B8, B53.

HXB2 Location Nef (72–91)

**Author Location** Nef (71–90 SF2)

Epitope PQVPLRPMTYKAAVDLSHFL

**Immunogen** HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

• CTL expanded ex vivo were later infused into HIV-1 infected patients.

HXB2 Location Nef (72-91)

**Author Location** Nef (SF2)

Epitope PQVPLRRMTYKAAVDLSHFL

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2001a

- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- Nef peptides PQVPLRRMTYKAAVDLSHFL, KAAVDLSH-FLKEKGGLEGLI and EEEEVGFPVTPQVPLRPMTY were recognized and the first two share KAAVDLSHFL (a Cw8 epitope), the first and last share PQVPLRPMTY.

**HXB2 Location** Nef (73–82)

**Author Location** Nef (73–82 NL43)

**Epitope** QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

References Koenig et al. 1990

- 81 Tyr is critical for binding to A3.1.
- C. Brander notes that this is an A\*0301 epitope in the 1999 database.

HXB2 Location Nef (73–82) **Author Location** Nef (73–82 LAI)

**Epitope QVPLRPMTYK** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords rate of progression, escape

References Koenig et al. 1995

- Alanine substitutions L76A, R77A, M79A, T80A significantly decreased immunogenicity of peptide.
- Nef CTL clones (4N225) were infused into an HIV-1 infected volunteer to evaluate effects of infusion on viral load/patient
- Infusion led to outburst of escape variants which resulted in higher viral load/accelerated disease progression.

HXB2 Location Nef (73-82)

**Author Location** Nef (73–82)

**Epitope** QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords immunodominance

References Betts et al. 2000

- to SLYNTVATL, calling into question whether it is immuno-
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals was A3, and responded to QV-PLRPMTYK as well as two other A3.1 epitopes.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82)

**Epitope** QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords acute/early infection References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects -3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.

No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

HXB2 Location Nef (73–82)
Author Location Nef (73–82 LAI)
Epitope QVPLRPMTYK
Subtype B
Immunogen
Species (MHC) human (A\*0301)

Keywords optimal epitope
References Frahm et al. 2007

• C. Brander notes this is an A\*0301 epitope.

HXB2 Location Nef (73–82) Author Location Nef (73–82) Epitope QVPLRPMTYK Subtype B

**Immunogen** in vitro stimulation or selection

Species (MHC) human (A\*0301)

Keywords epitope processing, dendritic cells

References Andrieu et al. 2003

- This study demonstrates that lipopeptides carrying epitopes can be taken up by human dendritic cells, processed using different pathways, and recognized by epitope-specific CD8+T-cells originally derived from HIV+ individuals. The RT ILKEPVHGV peptide was embedded in a longer peptide fragment in the lipopeptide, and was internalized by endocytosis and processed in the cytosol by proteasomal cleavage by following an endosome-to-cytosol pathway for processing and presentation. Administration of epoxomycin, a proteasome inhibitor, completely abrogated epitope presentation to a CD8+T-cell line, while monensin, an inhibitor of acid-dependent endosomal enzyme activity did not.
- In contrast to the RT epitope, dendritic cell presentation of the Nef epitope QVPLRPMTYK embedded in a longer peptide in a lipopeptide was not inhibited by epoxomycin, but was inhibited by monensin, indicative of endocytolitic epitope processing.

**HXB2 Location** Nef (73–82)

**Author Location** Nef

**Epitope** QVPLRPMTYK

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (A\*0301)

**Donor MHC** A\*0101, A\*0301, B\*0801, B\*5101

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFNγ, HLA binding

Keywords escape

References Milicic et al. 2005

 CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient. In an early acute infection of the donor, the precise timing of transmission determines the viral variants transmitted.

 Two variants of this epitope, QVPvRPMTYK and QaPLRPM-TYK, were found in 1 donor. The QaPLRPMTYK substitution reduced the binding affinity for A\*0301 by 52%.

HXB2 Location Nef (73–82)

**Author Location Nef** 

Epitope QVPLRPMTYK
Subtype A, B, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

**Species (MHC)** human, macaque (A\*0301, A11)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the A-clade epidemic in Nairobi, Kenya. A DNA and MVA prime-boost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location Nef (73–82)

Author Location (LAI)

**Epitope** QVPLRPMTYK

Subtype B

Immunogen

Species (MHC) human (A\*1101)

**Keywords** optimal epitope

References Buseyne 1999; Frahm et al. 2007

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82)

**Epitope** QVPLRPMTYK

Subtype B, CRF01\_AE

Immunogen

**Species (MHC)** (A\*1101)

Country Thailand.

Keywords HIV exposed persistently seronegative

(HEPS), immunodominance, structure

References Li & Bouvier 2004

• HLA-A\*1101 has been associated with resistance to acquisition of HIV-1 infection in female sex-workers in Thailand. Its crystal structure has been determined in association with two immunodominant A\*1101 HIV-1 CTL epitopes. Its anchor residues are confirmed as P2(Ile/Val) and C-term (Lys). The backbone conformation of the peptides is defined as two bulges separated by a secondary anchor residue (P6 Ser or Met) that may offer various advantages in the selection and presentation of CTL epitopes by HLA-A\*1101.

HXB2 Location Nef (73–82)
Author Location Nef (73–82)
Epitope QVPLRPMTYK
Immunogen HIV-1 infection
Species (MHC) human (A11)
References Le Borgne et al. 2000

 Soluble factors in supernatant from both an HIV-specific cloned CTL line and an EBV (Epstein-Barr-virus) CTL line inhibit viral replication, but do not block viral entry in CD4+ T lymphocytes, by a noncytotoxic mechanism.

HXB2 Location Nef (73–82)
Author Location Nef (73–82 LAI)
Epitope QVPLRPMTYK
Subtype B
Immunogen HIV-1 infection

**Species (MHC)** human (A11) **References** Robertson *et al.* 1993

Development of a retroviral vector (pNeoNef) to generate autologous CTL targets.

- Hunziker et al. [1998] suggests that HLA-A2 does not in fact present this epitope.
- The initial assignment of HLA-A2 presentation for this epitope was based on a serological HLA typing. Subsequently, the authors revisited the issue with genetic HLA typing and found that HLA-A11 was the correct presenting molecule (Dr. Florence Buseyne, pers. comm., 2000)

HXB2 Location Nef (73–82) Author Location Nef (73–82 LAI) Epitope QVPLRPMTYK Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

Keywords review, escape

References Couillin et al. 1994; Goulder et al. 1997a

- Mutational variation in HIV epitopes in individuals with appropriate HLA types can result in evasion of CTL response.
- Goulder *et al.* [1997a] is a review of immune escape that summarizes this study.

HXB2 Location Nef (73–82) Author Location Nef (73–82 LAI) Epitope QVPLRPMTYK Subtype B

Immunogen HIV-1 infection Species (MHC) human (A11) References Couillin *et al.* 1995

 Mutations found in this epitope in HLA-A11 positive and negative donors were characterized. HXB2 Location Nef (73–82)
Author Location Nef (73–82)
Epitope QVPLRPMTYK
Epitope name QVP
Immunogen HIV-1 infection

Species (MHC) human (A11)

Keywords HAART, ART, supervised treatment interruptions (STI), immunodominance, acute/early

infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- One of the 2/8 HLA-A11 study subjects recognized this CTL epitope.
- Patient SC18(HLA A2/11, B8/44, Cw06/0701, DR3/7, DR52/53, DQ2) recognizes the epitopes ACQGVGGPGHK, QVPLRPMTYK, AVDLSHFLK, and one called QIY but not fully described – he had brief therapy upon seroconversion and has had low viral load during 600 days of follow up.

**HXB2 Location** Nef (73–82) **Author Location** Nef (73–82)

Epitope QVPLRPMTYK

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

Author Location Nef (73–82)

Author Location Nef (73–82)

Epitope QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human (A11)

References Appay et al. 2000

- Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virus-specific CD8+ T cells produced IFN- $\gamma$  and MIP-1 $\beta$  with a distinct subset that failed to produce TNF- $\alpha$

HXB2 Location Nef (73–82)

Author Location Nef (71–80 93TH253 subtype CRF01)

**Epitope** QVPLRPMTYK **Epitope name** N73-82

Subtype CRF01\_AE

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was weakly reactive in HEPS study subjects 265 who was HLA A2/A11 and 128 who was HLA A11/A33, and after a second *in vitro* stimulation, in study subject 256 who was HLA A11/33, making it the most reactive epitope tested in HLA-A11 HEPS women, with 3/4 responding.
- This epitope was strongly reactive in HIV+ study subject 053 who carried HLA-A11.

HXB2 Location Nef (73-82)

**Author Location** Nef (71–80 93TH253 subtype CRF01)

Epitope QVPLRPMTYK
Subtype CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A11)

 $\textbf{Keywords} \ \ \text{subtype comparisons}$ 

References Bond et al. 2001

- HLA-A11 CRF01 (called subtype E in Bond *et al.*) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.
- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This epitope was predicted by the EpiMatrix method to be likely to bind to A11, and it served as an epitope in the FSWs, it was one of the six A11 epitopes that had been previously defined.
- 4/8 tested FSWs recognized this epitope.
- An HLA-A11 tetramer was made for this epitope, which was recognized by two subjects – only one subject had an expanded tetramer staining T-cell population after *in vitro* stimulation.
- This epitope was highly conserved in other subtypes, and exact matches were common.

HXB2 Location Nef (73–82)

**Author Location** Nef

Epitope QVPLRPMTYK

Epitope name QVP

**Immunogen** HIV-1 infection **Species (MHC)** human (A11)

Keywords HAART, ART, supervised treatment interrup-

tions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius et al. [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there
  was no correlation between CTL responses with viral rebound
  rates, plateau viral loads, or clearance rates.

HXB2 Location Nef (73–82)

**Author Location** Nef

**Epitope** QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human (A11)

Donor MHC A2, A11, B8, B60, Bw6

**Keywords** HAART, ART **References** Appay *et al.* 2002

- Four HIV patients with prolonged clinically successful antiviral therapy but with ongoing evidence of replication and Nef mRNA transcription, showed specific T-cell responses by Elispot and Tetramer staining, maintained for 2-4 years after initiation of HAART.
- Nef epitope recognition was detected in all 4 subjects, gp120,
   Pol and Gag-specific in 1 or 2 subjects two patients recognized this epitope, one using HLA-A3, one using HLA-A11.
- The HIV-specific CD8+ T-cells had an intermediate maturation phenotype characterized by low levels of perforin and high levels of CD27 expression.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82)

Epitope QVPLRPMTYK

Epitope name QK10

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (A11)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Flow cytometric T-cell cytokine

assay

Keywords optimal epitope

References Allen et al. 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- This epitope did not vary.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82)

Epitope QVPLRPMTYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

HXB2 Location Nef (73-82)

**Author Location Nef** 

**Epitope** QVPLRPMTYK

Epitope name QK9

**Immunogen** 

Species (MHC) (A11)

Keywords review, immunodominance, escape, acute/early infection, early-expressed proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

• This review discusses the importance of 3 factors that impact the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides listed as immunodominant in acute HIV-1 infection.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82)

Epitope QVPLRPMTYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11, A\*0301)

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, CD8 T-cell Elispot Author Location Nef (73–82 LAI) granzyme B

**Keywords** Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- None of three patients responded to this peptide with GzB producing cells, while all three responded with IFN-gamma producing cells.

HXB2 Location Nef (73-82)

**Author Location** Nef (73–81)

Epitope QVPLRPMTYK Immunogen HIV-1 infection

**Species (MHC)** human (A2, A3, A11, B35)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari et al. as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location Nef (73-82) **Author Location** Nef (73–82 LAI)

**Epitope** QVPLRPMTYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords epitope processing, escape

References Chassin et al. 1999

Mutations in Nef that flank this epitope, Thr71Lys and Ala83Gly, may account for an observed loss of CTL reactivity, with escape due to the introduction of proteasome processing defects.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82)

**Epitope** QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords subtype comparisons

References Durali et al. 1998

- Cross-clade CTL response was studied by determining the CTL activity in seven patients from Bangui, (6 A subtype, and 1 AG recombinant infections) and one A subtype infection from a person living in France originally from Togo, to different antigens expressed in vaccinia.
- Pol reactivity: 8/8 had CTL to A subtype, and 7/8 to B subtype, and HIV-2 Pol was not tested.
- Gag reactivity: 7/8 reacted with A or B subtype gag, 3/8 with HIV-2 Gag.
- Nef reactivity: 7/8 reacted with A subtype, and 5/8 with B subtype, none with HIV-2 Nef.
- Env reactivity: 3/8 reacted with A subtype, 1/8 with B subtype, none with HIV-2 Env.
- · One of the patients was shown to react to this epitope: QV-PLRPMTYK.

HXB2 Location Nef (73–82)

**Epitope** QVPLRPMTYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords review, escape

References Goulder et al. 1997e; Goulder et al. 1997a

- HLA-identical siblings, twin hemophiliac brothers, were both infected with the same batch of factor VIII.
- Both had a response to this epitope. One had a response to this epitope, the other did not.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82)

Epitope QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human (A3)

References Lubaki et al. 1997

- Eighty two HIV-1-specific CTL clones from 5 long-term nonprogressors were isolated and analyzed for breadth of response.
- · A sustained Gag, Env and Nef response was observed, and clones were restricted by multiple HLA epitopes, indicating a polyclonal response.

• An A3+ subject had a strong response to this epitope, with 10/11 CTL clones being specific for this epitope, isolated at two time points, 1 year apart.

HXB2 Location Nef (73–82) Author Location Nef (73–82) Epitope QVPLRPMTYK

Epitope name N1

Immunogen HIV-1 infection Species (MHC) human (A3)

**Keywords** HAART, ART, escape

References Samri et al. 2000

• The epitope was recognized by patients 252#0 and 252#4 in a study of the effects of therapy escape mutations on CTL recognition.

HXB2 Location Nef (73–82)
Author Location Nef (73–82 SF2)
Epitope QVPLRRMTYK
Immunogen HIV-1 infection
Species (MHC) human (A3)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A3+ individuals that had a CTL response to this epitope broken down by group: 3/7 group 1, 1/4 group 2, and 1/2 group 3.

HXB2 Location Nef (73–82)
Author Location Nef (SF2)
Epitope QVPLRPMTYK
Immunogen HIV-1 infection
Species (MHC) human (A3)
References Altfeld et al. 2000

 This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.

**HXB2 Location** Nef (73–82)

**Author Location** 

**Epitope** QVPLRPMTYK **Epitope name** Nef-QK10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3) References Sabbaj *et al.* 2003

 Among HIV+ individuals who carried HLA A03, 9/20 (45%) recognized this epitope. Author Location Nef (73–82)

Author Location Nef (73–82)

Epitope QVPLRPMTYK

Epitope name A3-QK10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3) Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 3/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 5/7 individuals had detectable responses to this epitope after STI.

HXB2 Location Nef (73-82)

**Author Location Nef** 

Epitope QVPLRPMTYK
Immunogen HIV-1 infection
Species (MHC) human (A3)
Donor MHC A3, B44, B64, Bw4, Bw6
Keywords HAAPT APT

**Keywords** HAART, ART **References** Appay *et al.* 2002

- Four HIV patients with prolonged clinically successful antiviral therapy but with ongoing evidence of replication and Nef mRNA transcription, showed specific T-cell responses by Elispot and Tetramer staining, maintained for 2-4 years after initiation of HAART.
- Nef epitope recognition was detected in all 4 subjects, gp120,
   Pol and Gag-specific in 1 or 2 subjects two patients recognized this epitope, one using HLA-A3, one using HLA-A11.
- The HIV-specific CD8+ T-cells had an intermediate maturation phenotype characterized by low levels of perforin and high levels of CD27 expression.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82)

**Epitope QVPLRPMTYK** 

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A3)

**Donor MHC** A1, A3, B7, B14, Cw\*0702, Cw\*0802

Assay type CD8 T-cell Elispot - IFNγ

Keywords binding affinity, acute/early infection, early-

expressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

HXB2 Location Nef (73-82) **Author Location** Nef (73–82) **Epitope** QVPLRPMTYK Immunogen HIV-1 infection Species (MHC) human (A3)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4, DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 17 potential epitopes from this individual, defined based on previously identified epitopes presented by appropriate HLA molecules. Full length genome sequence did not reveal changes in time in any of these epitopes over a four year period. Peptide pools initially revealed very little response to these eptiopes, but this increased over time.

HXB2 Location Nef (73-82)

**Author Location Nef** 

**Epitope** QVPLRPMTYK Immunogen HIV-1 infection Species (MHC) human (A3)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

Keywords HIV exposed persistently seronegative

(HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- 3/5 HLA A3+ infection-resistant men, compared to 1/3 preseroconversion men who went on to become infected, reacted to this epitope.

HXB2 Location Nef (73–82)

**Author Location** Nef (71–80)

**Epitope** QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human (A3)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong T-helper cell responses. Only patients starting with moderately high viral load (VL) were able to reduce the VL set point. HIV-1 specific CD8+ T-cell responses were shown to increase significantely until the end of the follow
- 9/14 patients recognized this epitope, it was the most recognized of six A\*03 epitopes.

HXB2 Location Nef (73–82)

Author Location (B consensus)

Epitope QVPLRPMTYK

Epitope name QK10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A03, B14, B60, Cw3, Cw7; A01, A03, B08, B14, Cw7, Cw8

Assay type Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.
- 2/9 individuals recognized this epitope, presented by HLA-A3.

HXB2 Location Nef (73-82)

**Author Location** Nef (73–82)

Epitope QVPLRPMTYK Immunogen HIV-1 infection Species (MHC) human (A3) **Donor MHC** A2, A3, B7, B44 Country United States.

> Assay type Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

> Keywords HAART, ART, escape, variant crossrecognition or cross-neutralization

References Casazza et al. 2005

- Evidence of continued viral evolution during HAART therapy despite low viral load was found in 1/5 patients studied.
- The epitope QVPLRPMTYK was invariant (18/18 sequences) prior to therapy in the patient that recognized it.

HXB2 Location Nef (73-82) **Author Location** Nef (73–82) **Epitope** QVPLRPMTYK Subtype C

Immunogen HIV-1 infection Species (MHC) human (A3)

**Donor MHC** A\*0301, A\*7401, B\*1510, B\*3501, Cw\*0401; A\*0301, A\*68, B\*0702, B\*1510, Cw\*0401, Cw\*0702

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection

References Pillay et al. 2005

- HIV-specific CTLs in infants were shown to be able to select for viral escape variants early in life, despite a lack of escape with the same CTL specificity in the mother. Infant CTL responses may be compromised by transmission of escape variants that arose in the mother and also those that arose in the father, if the father was the source of the mother's infection.
- QVPLRPMTYK is the C subtype consensus form of an epitope recognized in a mother, who carried this autologous variant: QVPLRPMTfK. QVPLRPMTfK was the dominant form in her infant at 2 weeks of age, but new variants rapidly emerged: QV-PLkPMTfK, QVPLRPMnYK, QVPvRPMTfK, QVPLRPMsYr, QVPLRPMsYK.

HXB2 Location Nef (73-82) **Author Location** Nef (73–82 LAI) **Epitope** QVPLRPMTYK

Epitope name N1

Subtype B

Immunogen HIV-1 infection **Species (MHC)** human (A3 supertype)

Keywords HAART, ART, supertype

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened – eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+

PBL – but with continued viral suppression, HIV-specific responses diminished.

• Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

HXB2 Location Nef (73–82) Author Location Nef (94-103) Epitope QVPLRPMTYK

Immunogen HIV-1 infection

**Species (MHC)** human (A3 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind 3/5 HLA-A3 supertype alleles (A\*0301, A\*1101, A\*3101, A\*3301 and A\*6801).

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82 BRU)

**Epitope** QVPLRPMTYK

Subtype B, CRF02 AG

Immunogen HIV-1 infection

Species (MHC) human (A3, A11)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02 AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 0/9 CRF02\_AG-infected Ivorians, and 2/9 B-infected French subjects.
- 3/8 Ivorians carried a substitution in this epitope, while only 1/5 B clade infected French people did, OVPvRPMTYK, and the substitution was found in one of the people that recognized the peptide.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82 BRU)

Epitope QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human (A3, A11, B35)

References Culmann et al. 1991

• Nef CTL clones from HIV+ donors.

HXB2 Location Nef (73–82) **Author Location** Nef (73–82 LAI) **Epitope** QVPLRPMTYK

Subtype B

Immunogen

Species (MHC) human (B27)

References Culmann 1998

• Optimal epitope mapped by peptide titration.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82 LAI)

**Epitope** SVPLRPMTYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35, Cw4)

**References** Buseyne *et al.* 1993a

- Vertical transmission of HIV ranges from 13% to 39%.
- Primary assays showed that cytotoxic activity against at least one HIV protein was detected in 70% of infected children.
- Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.
- Patient EM13, who had a CTL response to three epitopes in Nef, was infected via blood transfusion after birth and went from CDC stage P2A to P2E during the study.

HXB2 Location Nef (73–82)

**Author Location** 

Epitope QVPLRPMTYK

Epitope name OK10

Immunogen HIV-1 infection

Species (MHC) human (B8)

Country United States.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Tet-

ramer binding

Keywords immunodominance, acute/early infection,

characterizing CD8+ T cells, immune dys-

function

References Lichterfeld et al. 2004a

- HIV-1 specific CD8+ T-cells in acute and long-term nonprogressive HIV-1 infection show strong ex-vivo proliferative capacities which are rapidly lost in chronic HIV-1 infection. The loss of CD8+ T-cell function is closely linked with the loss of HIV-1 specific, IL2 secreting CD4+ T-cells. The function can be rescued in vitro and in vivo by restoring the specific CD4+ T-cell help.
- Full CD8+ T-cell responses to this epitope were dependent on co-stimulation with a CD4+ T cell dependent epitope from T-cells harvested during acute infection. The CD8+ T-cell response to this epitope was immunodominant in one study individual.

HXB2 Location Nef (73–82)

**Author Location** Nef (73–82)

Epitope QVPLRPMTYK

Immunogen HIV-1 infection

Species (MHC) human

References Garcia et al. 1997

- The anti-Nef CTL line P1 specific for this epitope is able to kill target cells via two mechanisms.
- First: Ca<sup>2+</sup>-dependent, perforin-dependent Nef-specific lysis.
- Second: Ca<sup>2+</sup>-independent, CD95-dependent apoptosis that could also kill non-specific targets.

- Findings indicate that the two mechanisms are not mutually exclusive in human CTL, as they are in mice.
- CTL mediated CD95-dependent apoptosis may play a role in pathogenesis.

HXB2 Location Nef (73–83)

**Author Location** Nef (73–82 BRU)

Epitope QVPLRPMTYKA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- QVPLRPMTYKA was recognized in 9/15 (60%) of individuals with HLA A3. It was a high affinity HLA-A3 binder.

**HXB2 Location** Nef (74–81)

Author Location Nef (74-82)

Epitope VPLRPMTY

Immunogen

Species (MHC) human (A3)

References Carreno et al. 1992

• Included in HLA-A3 binding peptide competition study.

HXB2 Location Nef (74–81)

**Author Location** Nef (73–82 LAI)

**Epitope** VPLRPMTY

Subtype B

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human (B\*3501)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*3501 epitope.

HXB2 Location Nef (74–81)

**Author Location** Nef (75–82)

**Epitope VPLRPMTY** 

Immunogen peptide-HLA interaction

Species (MHC) human (B\*3501)

References Smith et al. 1996

• Crystal structure of VPLRPMTY-class I B allele HLA-B\*3501 complex.

HXB2 Location Nef (74–81)

**Author Location** Nef

**Epitope VPLRPMTY** 

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

Keywords dendritic cells

References Ostrowski et al. 2000

- The role of CD4+ T-cell help in expansion of virus-specific memory CTL was studied through co-culture *ex vivo*
- Optimal expansion of HIV-1-specific memory CTL depended on CD4+ T cell help in 9 of 10 patients – CD40 ligand trimer (CD40LT) could enhance CTL in the absence of CD4+ T cell help to a variable degree in most of patients.
- Those CTL that didn't respond to CD40LT could expand with IL2 present, and IL15 produced by dendritic cells also contributes
- The T-helper epitope used for CD4+ T cell stimulation was the universal tetanus helper epitope TET830-843 (QYIKANSK-FIGITE)

HXB2 Location Nef (74–81)

Author Location Nef (74-81)

Epitope VPLRPMTY

Epitope name VY8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

 $\textbf{Donor MHC} \ A*0201, \ A*0301, \ B*3501, \ B*51, \ Cw*04,$ 

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, Chromium-release assay

Keywords escape, acute/early infection

References Bansal et al. 2005

- Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- Point mutation of epitope at position 8 (Y to F,VPLRPMTf) was detected at a chronic infection timepoint. The CTL response was strong in early infection, but diminished by month 13. This mutation had reduced avidity.

HXB2 Location Nef (74–81)

**Author Location** Nef (subtype B)

**Epitope** VPLRPMTY

Subtype B

**Immunogen** HIV-1 exposed seronegative

Species (MHC) human (B35)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix systemic CD8+ T cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

HXB2 Location Nef (74–81)

**Author Location** Nef

Epitope VPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords acute/early infection

References Wilson et al. 2000a

- Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B2705; and A\*0201, A\*0301, B2705, B39.
- ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects – 3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
- The subject with A\*0201 had a moderately strong response to SLYNTVATL.
- Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
- No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

HXB2 Location Nef (74–81)

Author Location Nef (73-82 LAI)

**Epitope** VPLRPMTY

Subtype B

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human (B35)

Keywords review

**References** Culmann *et al.* 1991; McMichael & Walker

 Review of HIV CTL epitopes – defined by B35 motif found within a larger peptide.

**HXB2 Location** Nef (74–81)

**Author Location** Nef (73–82 LAI)

**Epitope** VPLRPMTY

Subtype B

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human (B35)

References Rowland-Jones et al. 1995

VPLRPMTY also recognized by CTL from HIV-2 seropositives; epitope is conserved.

HXB2 Location Nef (74–81)

**Author Location** Nef

Epitope VPLRPMTY

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B35)

Keywords subtype comparisons, HIV exposed persis-

tently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades such cross-reactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A and D subtype consensus are identical to the B clade epitope.

HXB2 Location Nef (74–81) Author Location Nef (75–82) Epitope VPLRPMTY

Immunogen in vitro stimulation or selection

Species (MHC) human (B35)

References Lalvani et al. 1997

- A peptide-based protocol was optimized for restimulation of CTLp using optimized peptide and IL-7 concentrations importantly this protocol does not stimulate a primary response, only secondary peptide-specific CTLp counts could be obtained via staining with peptide-Class I tetramers.
- This peptide was one of the B35 presented test peptides used in control experiments showing that the assay gave no activity using lymphocytes from 21 healthy B35 seronegative donors.

HXB2 Location Nef (74–81) Author Location Nef (subtype B) Epitope VPLRPMTY

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B35)

Keywords subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found B clade epitopes are often cross-reactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among A, B, and D clade viruses.

HXB2 Location Nef (74–81)

**Author Location** Nef

Epitope VPLRPMTY

Immunogen

Species (MHC) human (B35)

References Rowland-Jones et al. 1999

- CTL responses in seronegative highly HIV-exposed African female sex workers in Gambia and Nairobi were studied these women had no delta 32 deletion in CCR5.
- In Gambia there is exposure to both HIV-1 and HIV-2, CTL responses to B35 epitopes in exposed, uninfected women are cross-reactive,
- HIV-2 version of this epitope is conserved: VPLRPMTY, and CTLs are cross-reactive one of five B35 CTL epitopes that are cross-reactive, see also Rowland-Jones *et al.* [1995]

**HXB2 Location** Nef (74–81) **Author Location** Nef (74–81)

**Epitope** VPLRPMTY

Epitope name VPL

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- One of two HLA B35+ among the eight study subjects recognized this epitope.
- Patient SC15 (HLA A1/68, B8/35, Bw4/6, Cw4/0704) was given acute and sustained therapy and recognized epitopes PPIPVGDIY and VPLRPMTY during 331 days of HAART treatment.

HXB2 Location Nef (74–81)

**Author Location** Nef (75–82)

**Epitope** VPLRPMTY

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B35)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Subject ML 857 shifted from a A\*6802 DTVLEDINL and B35 (H/N)PDIVIYQY response prior to seroconversion to a B35 PPIPVGDIY and B35 VPLRPMTY response postseroconversion.

HXB2 Location Nef (74–81)

**Author Location** 

Epitope VPLRPMTY

Epitope name Nef-VY8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Sabbaj et al. 2003

- Among HIV+ individuals who carried HLA B35, 12/22 (55%) recognized this epitope.
- Among HIV+ individuals who carried HLA B\*5301, 0/11 (0%) recognized this epitope.

HXB2 Location Nef (74–81)

**Author Location** Nef (74–81 BRU)

**Epitope** VPLRPMTY

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B35)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- VPLRPMTY was recognized in 5/16 (31%) of individuals with HLA B35, and it was a moderate affinity HLA binder. Cleavage at the C-term Y was frequent *in vitro*.

HXB2 Location Nef (74-81)

**Author Location** 

**Epitope** VPLRPMTY **Subtype** A, B, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade

HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (B35)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses
  to peptide pools were detected using intracellular cytokine
  staining and IFNgamma Elispot assays after vaccination of 5
  macaques. The response to the Mamu A\*01 SIV p27 epitope
  p11C (CTPYDINQM), included in the polyepitope region, was
  not immunodominant in the Mamu A\*01 vaccinated macaques,
  possibly because of processing limitations in context of the
  artificial polyepitope string Wee et al. [2002].

HXB2 Location Nef (74-81)

**Author Location** Nef (74–81)

**Epitope** VPLRPMTY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35)

**Donor MHC** A1, A3, B8, B35

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** acute/early infection, early treatment

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. SubjectBroadcast message from root Thu May 27 21:34:36 2004...n of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma vBattery Low Notification from APM BIOS (8% 0:12)or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

HXB2 Location Nef (74–81)

**Author Location** Nef (72–79)

Epitope VPLRPMTY

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 9 patients recognized this epitope.

HXB2 Location Nef (74–81)

**Author Location** Nef (C consensus)

Epitope VPLRPMTY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B35)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape

References Kiepiela et al. 2004

HLA class I restricted CD8+ T-cell responses against HIV-1
were analyzed in African patients. Significantly more responses
were shown to be HLA-B restricted. Viral load, CD4 count,
and thus rate of disease progression were also associated with
HLA-B alleles. In addition, the selection pressure imposed on

HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.

• People who carried B35 carried a variant of this epitope, while people who did not almost always carried the consensus form.

HXB2 Location Nef (74-81) **Author Location** Nef (72–79) Epitope VPLRPMTY Epitope name VPL Immunogen HIV-1 infection Species (MHC) human (B35)

**Donor MHC** A2, A68.1, B\*07, B\*3503, Cw\*0401, Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ

Keywords rate of progression, immunodominance, escape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant eptiopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- VPL epitope was one of six that were largely or completely replaced by escape variants, with the two escape forms coming up between days 172 and 635, vplrpmSy and vplrmptF.

HXB2 Location Nef (74–82) Author Location Nef (73–82) **Epitope** VPLRPMTYK

Immunogen peptide-HLA interaction

Species (MHC) human (A11) References Zhang et al. 1993

• Exploration of A11 binding motif.

HXB2 Location Nef (75–82) **Author Location** Nef (75–82 LAI) Epitope PLRPMTYK Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*1101)

Keywords review

References McMichael & Walker 1994

- Review of HIV CTL epitopes.
- C. Brander notes that this is an A\*1101 epitope in the 1999 database.

HXB2 Location Nef (75–82) **Author Location** Nef (75–82 LAI) **Epitope** PLRPMTYK Subtype B Immunogen HIV-1 infection Species (MHC) human (A\*1101)

Keywords optimal epitope References Frahm et al. 2007 • C. Brander notes this is an A\*1101 epitope.

HXB2 Location Nef (77-85)

**Author Location** Nef (77–85 LAI)

Epitope RPMTYKAAL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*0702)

Keywords escape

References Bauer et al. 1997

- Structural constraints on the Nef protein may prevent escape.
- Noted in Brander 1999, this database, to be B\*0702.

HXB2 Location Nef (77–85) **Author Location** Nef (77–85 LAI) **Epitope** RPMTYKAAL Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*0702)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is a B\*0702 epitope.

Author Location Nef (75–83 IIIB) Epitope RPMTYKAAL Immunogen HIV-1 infection Species (MHC) human (B7)

HXB2 Location Nef (77-85)

Keywords binding affinity, TCR usage References Oxenius et al. 2001b

- · Study of tetramer staining of B7 around RPMTYKAAL gave quantitative results that were very different than functional measurements based on an ELISPOT assay.
- Autologous clones were checked and 39/40 clones from two time points had the variant sequence RPMTYKGAL tetramers based on RPMTYKGAL gave a more intense and uniform staining and bound with higher affinity to the RPM-TYKGAL  $V\beta$ 14 TCR.

HXB2 Location Nef (77–85) **Author Location** Nef (77–85 SF2) Epitope RPMTYKAAL Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords HAART, ART, acute/early infection References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.

- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B7+ individuals that had a CTL response to this epitope broken down by group: 1/4 group 1, 0/3 group 2, and 1/1 group 3.

HXB2 Location Nef (77–85)
Author Location Nef (77–85)
Epitope RPMTYKAAL
Immunogen HIV-1 infection
Species (MHC) human (B7)

**Keywords** rate of progression, acute/early infection **References** Day *et al.* 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

HXB2 Location Nef (77–85) Author Location Nef (77–85) Epitope RPMTYKAAV Immunogen HIV-1 infection Species (MHC) human (B7)

**Keywords** rate of progression, acute/early infection **References** Day *et al.* 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

HXB2 Location Nef (77–85) Author Location Nef (77–85 BRU) Epitope RPMTYKAAV Subtype B Immunogen HIV-1 infection Species (MHC) human (B7)

**Keywords** binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- RPMTYKAAV was recognized in 7/10 (70%) of individuals with HLA B7, and 0/3 (0%) of individuals with HLA B35. It was a moderate affinity HLA binder.

HXB2 Location Nef (77–85) Author Location Nef (77–85) Epitope RPMTYKAAL

Epitope name B7-RL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 3/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 2/4 individuals had detectable responses to this epitope after STI.

HXB2 Location Nef (77–85)

**Author Location** Nef (77–85)

Epitope RPMTYKAAV

Epitope name B7-RV9

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

 CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.

- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 2/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection - 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 3/4 individuals had detectable responses to this epitope after STI.

HXB2 Location Nef (77-85)

Author Location Nef (75-83)

Epitope RPMTYKAAL Immunogen HIV-1 infection Species (MHC) human (B7)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 3/7 patients recognized this epitope.

HXB2 Location Nef (77–85)

**Author Location** Nef (75–83)

Epitope RPMTYKGAL

**Epitope name** RPM

Immunogen HIV-1 infection Species (MHC) human (B7)

Cw\*0702, DR17, DR15, DR51, DR52, DQ2,

D<sub>0</sub>6

Assay type CD8 T-cell Elispot - IFNγ, Tetramer binding, CD4 T-cell Elispot - IFNγ

Keywords rate of progression, immunodominance, escape

References Oxenius et al. 2004b

- The increase in plasma viral load in a patient that progressed rapidly was preceded by positive selection of viral escape mutations in epitopes targeted by dominant HIV-1-specific CD8+ T-cell responses, and a decrease in HIV-1-specific CD4+ and CD8+ T-cell frequencies. Overall, escape variant epitopes were recognized 0-80% as efficiently as the index peptide, and the relatively efficiency of the variant eptiopes increased using PBLs collected after their appearance. No changes were found in viral tropism, replication kinetics and neutralizing antibody titers, so the rapid decline of the patient was attributed to loss of HIV containment due to CTL escape.
- This epitope was one of six epitopes found to be under positive selection for escape mutations and was completely replaced by

escape variants between days 172 and 635 (rpmtFkgal, rpm-SvkAal, rpmtvkgaV, rpmtvkAal). The first two were the most common at day 635, and experimentally shown to be escape.

HXB2 Location Nef (77–85)

**Author Location Nef** 

Epitope RPMTYKGAL

Epitope name RL9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- · Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 4, RPMnYKGAL, was found not to correspond to the most polymorphic residue in the epitope.

HXB2 Location Nef (77–85)

**Author Location** Nef (77–85 BRU)

**Epitope** RPMTYKAAV

Subtype B, CRF02 AG

Immunogen HIV-1 infection

Country Cote D'Ivoire.

Species (MHC) human (B7)

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons

References Inwoley et al. 2005

- Donor MHC A2, A68.1, B\*07, B\*3503, Cw\*0401, CD8+ T-cells from HIV-1 CRF02 AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
  - This epitope was recognized by 0/9 CRF02\_AG-infected Ivorians, and 2/9 B-infected French subjects.
  - One of the B-clade infected subjects that recognized this peptide was not sequenced, the other had one amino acid change: RPMTYKAAl. A variant form was in 3/5 B clade infection sequences. 8/8 CRF01 infected individuals had a variant of this peptide.

HXB2 Location Nef (77–85)

**Author Location** Nef (77–85)

Epitope RPMTYKAAL

Epitope name RW9

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A\*24, A\*30, B\*39, B\*07, Cw\*12, Cw\*17

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

mission, escape, characterizing CD8+ T cells, reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs. Escape variants were commonly detected in maternal plasma. Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- The variant RPMThqAAw was present in 10/10 clones from a B7- mother, was transmitted to her B7+ infant, and present in 29/30 clones at months 2, 6, and 152.

HXB2 Location Nef (77-85) Author Location Nef (79-85)

**Epitope** RPMTYKAAV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A3, A33, B14, B35, Cw\*0401, Cw\*0802

Assay type CD8 T-cell Elispot - IFNγ

Keywords acute/early infection, early treatment

References Cao et al. 2003

- · All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized; 24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- · More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

HXB2 Location Nef (77–91)

**Author Location** Nef (77–91)

Epitope RPMTYKGAFDLSFFL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons References Novitsky et al. 2002

- Keywords responses in children, mother-to-infant trans• HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
  - Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
  - This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Nef (79–87)

Author Location Nef (81–89 HXB3)

**Epitope** MTYKAALDL

Immunogen vaccine

Vector/Type: DNA, peptide Strain: B clade HXB3 HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (A\*0201)

Keywords binding affinity, computational epitope predic-

tion

References Sandberg et al. 2000

- Ten Nef 9-mer peptides were predicted to have a strong binding affinity with HLA-A\*0201 – of these, four did bind strongly by a T2 class I stabilization assay, several others bound weakly.
- A CTL immune response to only 3/10 peptides was detected by a 51Cr-release assay after immunization of HLA-A201 transgenic mice with either nef DNA under the control of a CMV promotor coated on, gold particles delivered to abdominal skin by gene gun.
- MTYKAALDL bound weakly to HLA-A2, but the DNA nef vaccine elicited a good CTL response.

HXB2 Location Nef (79–87)

**Author Location** Nef (79–87)

Epitope MTYKAALDL

Epitope name Nef79-87

Immunogen HIV-1 infection

Species (MHC) human, humanized mouse (A2)

Country United States.

Assav type CD8 T-cell Elispot - IFNγ

Keywords responses in children, immunodominance,

characterizing CD8+ T cells

References Chandwani et al. 2004

- Responses to five HLA-A2 presented epitopes in Nef were characterized in a population of 19 HIV infected adults and 21 children. The CD8 T-cell response to Nef was stronger and broader in adults than children, averaging 652 and 87 SFCs/10<sup>6</sup> PBMC, and 2.6 and 0.9 recognized epitopes per person, respectively.
- This is one of three novel Nef epitopes previously identified in HLA-A2 transgenic mice, shown to induce CD8 T-cell response in humans. It was not the immunodominant response.

HXB2 Location Nef (79-87)

**Author Location** Nef

Epitope MTYKAAVDL

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human (B63)

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, cross-presentation by dif-

ferent HLA

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This is a putative HLA-B63/57/58 epitope containing the B58 supertype binding motif. There is no evidence for B57/B58 cross-presentation of this epitope.

HXB2 Location Nef (79–93)

**Author Location** Nef

**Epitope MTYKGAFDLSHFLKE** 

Subtype A, D

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*6601, A\*6801, B\*5301, B\*5802; A\*0202,

A\*3002, B\*5703, B\*5802

Country Uganda.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, characterizing CD8+ T

cells

References Barugahare et al. 2005

- T-cell immune responses were examined in HIV-1 infected Ugandans. Similar levels of cross-clade responses for Gag, Env and Nef were observed. Higher frequencies of responses in conserved regions were found for Gag within a single clade, while areas with higher viral sequence variation had fewer detectable responses. Differential interclade sequence homologies between the Gag regions did not affect the respective level of cross-clade recognition.
- Novel unmapped epitope, this test peptide was conserved in the people that recognized it.

HXB2 Location Nef (80–87)

Author Location Nef (80-87)

**Epitope** TYKAAVDL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (A24)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

Keywords vaccine-induced epitopes, variant cross-

recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

HXB2 Location Nef (80–87)

Author Location Nef (80–87 BRU)

**Epitope** TYKAAVDL

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (A24)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 1/9 CRF02\_AG-infected Ivorians, and 0/9 B-infected French subjects.
- This epitope was highly variable in Ivorians; 8/9 had amino acid substitutions. The only one that reacted carried the sequence TYKgAfDL. 3/5 of the B clade French subjects carried variants, which tended to have only 1 amino acid substitution.

HXB2 Location Nef (80–94)

Author Location Nef (80–94 HXB2)

**Epitope** TYKAAVDLSHFLKEK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 47% of the study subjects, and it was the most frequently recognized peptide.

HXB2 Location Nef (82–90)

Author Location (C consensus)

Epitope KGAFDLSFF

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*57)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location Nef (82–90)

**Author Location** Nef (82–90)

Epitope KAAVDLSHF

Epitope name KF9

Subtype B, C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*57, B\*5801)

Country Australia.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, HLA binding

Keywords subtype comparisons, computational epitope

prediction, mother-to-infant transmission, escape, reversion, viral fitness, optimal epitope

References Leslie et al. 2005

• KAAVDLSHF is the susceptible optimal form of the epitope, and KgAVDLSHF an escape variant. The KgAVDLSHF form of the epitope was shown to be an escape mutation by virtue of an increased off-rate; however Elispot reactions to both forms are positive. The escape form was shown to be transmitted, and the most common form of the epitope in a B clade infected population in Perth, Australia (52%).

HXB2 Location Nef (82–90)

Author Location Nef (C consensus)

**Epitope KAAFDLSFF** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, reversion, viral fitness

References Kiepiela et al. 2004

HLA class I restricted CD8+ T-cell responses against HIV-1
were analyzed in African patients. Significantly more responses
were shown to be HLA-B restricted. Viral load, CD4 count,
and thus rate of disease progression were also associated with
HLA-B alleles. In addition, the selection pressure imposed on
HIV-1 by HLA-B alleles was shown to be substantially greater
than by other alleles.

 People who carried B57 all carried a variant of this epitope, while about half of the people who did not carry B57 carried the susceptible form, suggesting there is not a high fitness cost and revision rate in this case.

• HLA-B57 was associated with a low viral load.

HXB2 Location Nef (82–90)

Author Location Nef (82–90)

Epitope KAAFDLSFF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

**Donor MHC** A\*3001, A\*66, B\*4201, B\*5802, Cw\*0602,

Cw\*1701; A\*68, A\*66, B\*57, B\*5802,

Cw\*0602, Cw\*0701

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords responses in children, mother-to-infant trans-

mission, escape, acute/early infection

References Pillay et al. 2005

- HIV-specific CTLs in infants were shown to be able to select for viral escape variants early in life, despite a lack of escape with the same CTL specificity in the mother. Infant CTL responses may be compromised by transmission of escape variants that arose in the mother and also those that arose in the father, if the father was the source of the mother's infection.
- KAAFDLSFF is the C consensus form of the epitope; the autologous form in the mother was KAAFDLgFF, and this was transmitted to her infant. By 33 weeks a new dominant form of the epitope had emerged in the infant: gAAFDLgFF.

HXB2 Location Nef (82–91)

**Author Location** Nef (82–91)

**Epitope** KAALDLSHFL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade

LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (A2)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

Keywords vaccine-induced epitopes, variant cross-

recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

- After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of 31 that the vaccinated volunteers responded to.
- A KAAvDLSHFL variant was cross-recognized after the last boost.

HXB2 Location Nef (82–91)

**Author Location** Nef (82–91 LAI)

Epitope KAAVDLSHFL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (Cw\*0802) Keywords HAART, ART References Nixon *et al.* 1999

- A patient who made a mono-specific CTL response to this Nef specific epitope was given effective anti-retroviral therapy within 90 days of infection, reducing the antigenic stimulous.
- Within 7 days of therapy, his CTLp frequency dropped from 60 to 4 per million PBMC, as his viremia dropped.
- The patient went from having an activated effector population (detected by CTLp and clone specific RNA) to a nonactivated quiescent population (detected by the CTL-clone specific DNA)

Author Location Nef (82–91)
Author Location Nef (82–91 SF2)
Epitope KAAVDLSHFL
Immunogen HIV-1 infection
Species (MHC) human (Cw8)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-Cw8+ individuals that had a CTL response to this epitope broken down by group: 1/3 group 1, 0/0 group 2, and 0/1 group 3.

HXB2 Location Nef (82–91)
Author Location Nef (SF2)
Epitope KAAVDLSHFL
Immunogen HIV-1 infection
Species (MHC) human (Cw8)
References Altfeld et al. 2000

• This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.

HXB2 Location Nef (82–91)
Author Location (B consensus)
Epitope KAAVDLSHFL
Epitope name KL10
Subtype B
Immunogen HIV-1 infection

Species (MHC) human (Cw8)

 $\textbf{Donor MHC}\ \ A25, A32, B08, B14, Cw7, Cw8$ 

Country United States.

Assay type Cytokine production, Intracellular cytokine staining, Chromium-release assay, Flow cytometric T-cell cytokine assay **Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

HXB2 Location Nef (82–91)

**Author Location** Nef

Epitope KAAVDLSHFL

Epitope name KL10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (Cw8)

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 6, KAAVDmSHFL, was found not to correspond to the most polymorphic residue in the epitope.

HXB2 Location Nef (82–91)

**Author Location** 

Epitope KAAVDLSHFL

Epitope name KL10

Immunogen

Species (MHC) human (Cw8)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a Cw08 epitope.

HXB2 Location Nef (82–96)

**Author Location** Nef (82–96)

Epitope KGAFDLSFFLKEKGG

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.

• This peptide was among the 8 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** Nef (82–101)

Author Location Nef (81-100 SF2)

Epitope KAAVDLSHFLKEKGGLEGLI

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- Three of these 11 had CTL response to this peptide.
- The responding subjects were HLA-A1, A2, B8, B14; HLA-A11, A24, B8, B53.

HXB2 Location Nef (82–101)

**Author Location** Nef (SF2)

Epitope KAAVDLSHFLKEKGGLEGLI

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2001a

- HIV+ individual AC-06 was tested for reactive overlapping peptides spanning all HIV-1 proteins in an ELISPOT and was found to react with 12 peptides from 7 proteins, suggesting that the breadth of CTL responses are underestimated if accessory proteins are not included in the study.
- Nef peptides PQVPLRRMTYKAAVDLSHFL, KAAVDLSH-FLKEKGGLEGLI and EEEEVGFPVTPQVPLRPMTY were recognized and the first two share KAAVDLSHFL (a Cw8 epitope), the first and last share PQVPLRPMTY.

HXB2 Location Nef (83–90)

Author Location Nef (83–90 HXB2)

**Epitope** AAVDLSHF

Subtype B, CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (B62)

Country Viet Nam.

Assay type HLA binding

**Keywords** subtype comparisons, computational epitope prediction, escape, variant cross-recognition or cross-neutralization, vaccine antigen de-

sign

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-All, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- CRF01\_AE variant GaFdlsFf had a higher HLA-B62 binding score than the HXB2 epitope.

HXB2 Location Nef (83–91)

**Author Location** Nef (85–93 HXB3)

**Epitope** AALDLSHFL

Immunogen vaccine

Vector/Type: DNA, peptide Strain: B clade HXB3 HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (A\*0201)

**Keywords** binding affinity, computational epitope prediction

References Sandberg et al. 2000

- Ten Nef 9-mer peptides were predicted to have strong binding affinity for HLA-A\*0201 – of these, four did bind strongly by a T2 class I stabilization assay, several others bound weakly.
- A CTL immune response to only 3/10 peptides was detected by a 51Cr-release assay after immunization of HLA-A201 transgenic mice with nef DNA under the control of a CMV promotor, coated on gold particles delivered to abdominal skin by gene gun.
- AALDLSHFL was predicted to have a strong binding capacity for HLA-A2, and did, but it was the only one of the peptides recognized that was a strong binder, the other two recognized peptides were weak binders.
- AALDLSHFL was also tested by subcutaneous injection of Nef peptides in Freund's adjuvant and gave a strong response to the peptide.

HXB2 Location Nef (83–91)

Author Location (C consensus)

**Epitope** GAFDLSFFL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*0205)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- GAFDLSFFL is an optimal epitope.

**HXB2 Location** Nef (83–91)

**Author Location** Nef (83–91)

Epitope GAFDLSFFL

Epitope name GL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0205)

Keywords optimal epitope

References Frahm et al. 2007

C. Brander notes this is a A\*0205 epitope.

HXB2 Location Nef (83–91)

**Author Location** Nef (83–91 BRU)

Epitope AAVDLSHFL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- AAVDLSHFL was recognized in 3/18 (17%) of individuals with HLA A2. It was a low affinity HLA binder.

HXB2 Location Nef (83–91)

**Author Location** Nef (83–91)

Epitope AAVDLSHFL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (A2)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

Keywords vaccine-induced epitopes, variant cross-

recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of 31 that the vaccinated volunteers responded to.

HXB2 Location Nef (83–91)

Author Location Nef (83-91)

Epitope AALDLSHFL

Epitope name Nef83-91

Immunogen HIV-1 infection

Species (MHC) human, humanized mouse (A2)

Country United States.

Assav type CD8 T-cell Elispot - IFNγ

Keywords responses in children, immunodominance,

characterizing CD8+ T cells

References Chandwani et al. 2004

- Responses to five HLA-A2 presented epitopes in Nef were characterized in a population of 19 HIV infected adults and 21 children. The CD8 T-cell response to Nef was stronger and broader in adults than children, averaging 652 and 87 SFCs/10<sup>6</sup> PBMC, and 2.6 and 0.9 recognized epitopes per person, respectively.
- The novel AALDLSHFL Nef epitope was the most frequently and most strongly recognized epitope in this study, making it a possible immunodominant epitope.
- This is one of three novel Nef epitopes previously identified in HLA-A2 transgenic mice, shown to induce CD8 T-cell response in humans.

HXB2 Location Nef (83–91)

Author Location Nef (83–91 BRU)

**Epitope** AAVDLSHFL

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

**Species (MHC)** human (A2)

Country Cote D'Ivoire.

Assav type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons

References Inwolev et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 1/9 CRF02\_AG-infected patients, and by 1/9 B-infected patients. Variants were present in 6/8 Ivorians, and in 3/5 French subjects.
- The Ivorian who recognized the B clade peptide carried the substitutions gAfDLSHFL.

HXB2 Location Nef (83–91)

**Author Location** Nef (83–91 HXB2)

Epitope AAVDLSHFL

Subtype B, CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country Viet Nam.

Assay type HLA binding

**Keywords** subtype comparisons, computational epitope prediction, escape, variant cross-recognition or cross-neutralization, vaccine antigen de-

sign

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-A11, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- The CRF01\_AE variant GaFdlsFfl had a higher HLA-A2 binding score than the HXB2 epitope.

HXB2 Location Nef (83–91)

**Author Location** Nef (83–91)

Epitope AAVDLSHFL

Epitope name AL9

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A\*02, A\*30, B\*18, B\*13, Cw\*01, Cw\*05; A\*02, A\*32, B\*07, B\*40, Cw\*03, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords responses in children, mother-to-infant trans-

mission, escape, characterizing CD8+ T cells,

reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs. Escape variants were commonly detected in maternal plasma. Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- An escape form of the A2 epitope, AAVDmSHFL, was transmitted from an A2- mother to her A2+ infant, where it persisted in 29/29 sequences sampled over 11 months.
- Another form of this A2 epitope, gAlDLSHFL, was transmitted by an A2+ mother to an A2- infant, where it persisted in 30/30 sequences sampled over 15 months.
- AAVDmSHFL was shown to have lower responder cell frequencies than AAVDLSHFL.

HXB2 Location Nef (83–91) Author Location Nef (83–91) Epitope AAVDLSHFL Immunogen HIV-1 infection

Species (MHC) human (B60, B62, Cw\*0802, Cw8)

**Donor MHC** A\*0201, A23, B44, B62, Cw3, Cw4; A1, A3, B7, B14, Cw\*0702, Cw\*0802; A\*0201, A31, B44, B60, Cw3, Cw16; A1, A1, B8, B14, Cw7, Cw8

Assay type CD8 T-cell Elispot - IFNγ

Keywords acute/early infection, early treatment

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- Four different individuals recognized this epitope during a primary infection, and it was shown to be presented by HLA B60, B62, C2\*0802, and Cw8.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

HXB2 Location Nef (83–91) Author Location Nef (83–91 LAI) Epitope AAVDLSHFL

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Subtype B
Immunogen HIV-1 infection
Species (MHC) human (Cw*0802)
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**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a C\*0802(Cw8) epitope.

HXB2 Location Nef (83-91)

**Author Location** 

Epitope GAFDLSFFL

Epitope name GL9

**Immunogen** 

Species (MHC) human (Cw\*0802)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a Cw\*0802 epitope.

HXB2 Location Nef (83–91)

**Author Location** Nef (83–91)

Epitope AALDLSHFL

Immunogen

Species (MHC) human (Cw3)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

HXB2 Location Nef (83–92)

**Author Location** Nef (81–90 93TH253 subtype CRF01)

**Epitope** GAFDLSFFLK

Epitope name N83-92

Subtype CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Sriwanthana et al. 2001

- This was a study of HIV-1 exposed persistently seronegative (HEPS) female sex workers in Chiang Mai, northern Thailand.
- HLA-A11 is very common in this population, and was enriched among the HEPS sexworkers – weak CTL responses were detected in 4/7 HEPS women, and CTL responses were found in 8/8 HIV+ controls, and 0/9 HIV- women that were not exposed.
- This epitope was strongly reactive in HIV+ study subjects 053 and 184 who carried HLA-A11.

HXB2 Location Nef (83–92)

**Author Location** Nef (81–90 93TH253 subtype CRF01)

Epitope GAFDLSFFLK
Subtype CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A11)

**Keywords** subtype comparisons **References** Bond *et al.* 2001

 HLA-A11 CRF01 (called subtype E in Bond et al.) epitopes were identified that stimulated CTL from HIV+ female sex workers (FSW) from Northern Thailand, of whom more than half were HLA-A11 positive.

- 77 possible HLA-A11 epitopes were first defined using Epi-Matrix, these were screened for binding to A11 finding and 26 bound, and 12 of these were epitopes for CTL responses from 8 HLA-A11 positive FSWs, six were novel, six were previously identified.
- This epitope was predicted by the EpiMatrix method to be likely to bind to A11, and it served as an epitope in the FSWs, it was one of the six A11 epitopes that had been previously defined.
- 4/8 tested FSWs recognized this epitope.
- This epitope was only conserved in CRF01 and subtype C, and exact matches were uncommon.

HXB2 Location Nef (83–92) Author Location Nef (83–92) Epitope AAVDLSHFLK Subtype B

Immunogen HIV-1 infection Species (MHC) human (A11)

**Donor MHC** A\*0201, A11, B51, B61, Cw2, Cw\*14

Assay type CD8 T-cell Elispot - IFNγ

Keywords acute/early infection, early treatment

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.

HXB2 Location Nef (83–94)
Author Location Nef (83–94 BRU)
Epitope AAVDLSHFLKEK
Immunogen HIV-1 infection
Species (MHC) human (A11)
References Culmann et al. 1991

 Epitope defined by boundaries of overlapping peptides that stimulate Nef CTL clones.

HXB2 Location Nef (84–91) Author Location Nef (84–91) Epitope AVDLSHFL Subtype B Immunogen vaccine Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: OS21

Species (MHC) human (A2)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of 31 that the vaccinated volunteers responded to.

HXB2 Location Nef (84–91)
Author Location Nef (84–91 LAI)
Epitope AVDLSHFL
Subtype B
Immunogen HIV-1 infection
Species (MHC) human (B62)

References Culmann-Penciolelli et al. 1994

HXB2 Location Nef (84–91)
Author Location Nef (84–91)
Epitope AVDLSHFL
Immunogen HIV-1 infection
Species (MHC) human (B62)
Keywords immunodominance
References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals that didn't respond to SLYNTVATL reacted with seven other epitopes including this epitope.

Author Location Nef (84–91)
Author Location Nef (84–91 BRU)
Epitope AVDLSHFL
Subtype B, CRF02\_AG
Immunogen HIV-1 infection
Species (MHC) human (B62)
Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ
Keywords subtype comparisons

**References** Inwoley *et al.* 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 2/9 CRF02\_AG-infected Ivorians, and 0/9 B-infected French subjects.

• The 2 CRF02 infected subjects that recognized this peptide carried a form with one amino acid change: AfDLSHFL, Variant forms were in 4/8 CRF02 infection sequences. 3/5 B clade infected individuals had a variant of this peptide.

HXB2 Location Nef (84-92) Author Location Nef (84–92) Epitope AVDLSHFLK Immunogen HIV-1 infection Species (MHC) human (A\*0301) Keywords optimal epitope References Frahm et al. 2007

HXB2 Location Nef (84–92) Author Location Nef (84–92 LAI) **Epitope** AVDLSHFLK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A\*1101) **Keywords** optimal epitope References Frahm et al. 2007

• C. Brander notes this is an A\*1101 epitope.

HXB2 Location Nef (84–92) **Author Location** Nef (84–92) **Epitope** AVDLSHFLK Subtype B, CRF01\_AE Immunogen HIV-1 infection Species (MHC) human (A\*1101) **Keywords** subtype comparisons References Fukada et al. 2002

- Counterparts for eight known clade B HLA A\*1101 epitopes were generated for clade E (CRF01). Three epitopes, identical among clade A-E, were cross-reactive and recognized by clade E infected individuals. The clade E and B analogs to three more HLA A\*1101 epitopes was recognized in a clade specific manner. Two other HLA A\*1101 clade B defined epitopes were found not to have stimulated a response in clade E infected individuals.
- · AVDLSHFLK was found to elicit clade-specific responses in clade B (AVDLSHFLK is most common, aLdlshflk is a common variant also found in clade A) and clade E (aFdlsFflk is most common and is also common in clade C). AVDLSHFLK was strongly recognized by CTL from 2/5 B clade infected Japanese subjects, as was aLdlshflk, and aFdlsFflk by CTL from 5/7 E clade infected Thai subjects.
- The binding of aFdlsFflk to HLA A\*1101 was 10-50 times lower than the other variants, and bulk CTL generated from individuals did not cross-react with the cross-clade peptides.

HXB2 Location Nef (84–92) **Author Location** Nef (84–92 LAI) **Epitope** AVDLSHFLK Subtype B Immunogen HIV-1 infection Species (MHC) human (A11)

Kevwords review

References McMichael & Walker 1994

• Review of HIV CTL epitopes.

• C. Brander notes that this is an A\*1101 epitope in the 1999 database.

HXB2 Location Nef (84–92) Author Location Nef (84-92) Epitope AVDLSHFLK Immunogen HIV-1 infection Species (MHC) human (A11) Keywords immunodominance References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immuno-
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals that didn't respond to SLYNTVATL reacted with seven other epitopes including this epitope.

HXB2 Location Nef (84–92) **Author Location** Nef (84–92 LAI) **Epitope** AVDLSHFLK Subtype B Immunogen HIV-1 infection Species (MHC) human (A11) Keywords review, escape

References Couillin et al. 1994; Goulder et al. 1997a

- · Mutational variation in HIV epitopes in individuals with appropriate HLA types can result in evasion of CTL response.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study.

Author Location Nef (84-92 LAI) **Epitope** AVDLSHFLK Subtype B Immunogen HIV-1 infection Species (MHC) human (A11) References Couillin et al. 1995

HXB2 Location Nef (84–92)

• Mutations found in this epitope in HLA-A11 positive and negative donors were characterized.

HXB2 Location Nef (84–92) **Author Location** Nef (84–92) Epitope AVDLSHFLK Epitope name AVD **Immunogen** HIV-1 infection Species (MHC) human (A11)

> Keywords HAART, ART, supervised treatment interruptions (STI), immunodominance, acute/early infection

References Oxenius et al. 2000

• Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.

- Both of the 2/8 HLA-A11 study subjects recognized this CTL epitope.
- Patient SC19(HLA A11/12, B8/44, Cw06/0701, DR3/7, DR52/53, DQ 2/8) had a CTL response to epitopes FLKEKGGL, GEIYKRWII, ACQGVGGPGHK, AVDLSH-FLK, and FNCGGEFFY that declined during therapy initiated at day 197.
- Patient SC18(HLA A2/11, B8/44, Cw06/0701, DR3/7, DR52/53, DQ2) recognizes the epitopes ACQGVGGPGHK, QVPLRPMTYK, AVDLSHFLK, and one called QIY but not fully described – he had brief therapy upon seroconversion and has had low viral load during 600 days of follow up.

Author Location Nef (84–92)
Author Location Nef (82–90)
Epitope AVDLSHFLK
Immunogen HIV-1 infection
Species (MHC) human (A11)
References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location Nef (84–92)
Author Location Nef (84–92 SF2)
Epitope AVDLSHFLK
Immunogen HIV-1 infection
Species (MHC) human (A11)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A11+ individuals that had a CTL response to this epitope broken down by group: 0/3 group 1, 0/0 group 2, and 2/2 group 3.

HXB2 Location Nef (84–92) Author Location Nef (84–92) Epitope AVDLSHFLK

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC) human (A11)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers. HXB2 Location Nef (84–92)

**Author Location** Nef

**Epitope** AVDLSHFLK

Epitope name AVD

Immunogen HIV-1 infection Species (MHC) human (A11)

**Keywords** HAART, ART, supervised treatment interruptions (STI)

tions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius *et al.* [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there was no correlation between CTL responses with viral rebound rates, plateau viral loads, or clearance rates.

HXB2 Location Nef (84–92)

**Author Location Nef** 

Epitope AVDLSHFLK

**Subtype** A, B, D, F

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (A11)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location Nef (84–92)

Author Location Nef (84-92)

Epitope AVDLSHFLK

Epitope name AK9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A11)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining. Flow cytometric T-cell cytokine

assay

**Keywords** optimal epitope **References** Allen *et al.* 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- This epitope did not vary.

HXB2 Location Nef (84–92)

**Author Location** Nef (84–92)

**Epitope** AVDLSHFLK

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A11)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

HXB2 Location Nef (84–92)

Author Location Nef (B consensus)

Epitope AVDLSHFLK

Epitope name AK9

Immunogen HIV-1 infection Species (MHC) human (A11, A3)

**Donor MHC** A02, A11, B18, B44, Cw5, Cw12; A03, B14,

B60, Cw3, Cw7

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

**Keywords** assay standardization/improvement, crosspresentation by different HLA, memory cells,

characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 2/9 individuals recognized this epitope, each in the context of a different HLA-presenting molecule.

HXB2 Location Nef (84–92)

Author Location Nef (84-92 BRU)

**Epitope** AVDLSHFLK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Keywords** binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- AVDLSHFLK was recognized in 4/12 (33%) of individuals with HLA A3. It was a high affinity HLA-A3 binder.

HXB2 Location Nef (84–92)

**Author Location** Nef (84–94)

Epitope AVDLSHFLK

**Epitope name** A3-ALK9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/14 HLA-A3 positive individuals had detectable A3-restricted responses to this epitope during acute infection, but only 5/15 of HLA-A3 epitopes tested were targeted during acute infection. 1/7 individuals began to have detectable responses to this epitope after STI.

HXB2 Location Nef (84–92)

**Author Location** Nef (84–92)

Epitope AVDLSHFLK

**Epitope name** AK9

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Donor MHC** A\*03, A\*31, B\*08, B\*15, Cw\*04, Cw\*07; A\*24, A\*31, B\*47, B\*15, Cw\*04, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, characterizing CD8+ T cells,

reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs.
   Escape variants were commonly detected in maternal plasma.
   Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- Variant sequence aMdlshflk was present in 7/10 clones from A3+ mother, was transmitted and present in 10/10 clones at months 2 and 4, but dropped to 0/10 clones by 15 months of age in her A3- child.

HXB2 Location Nef (84–92)

**Author Location** Nef (84–92)

**Epitope** AVDLSHFLK

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (A3, A11)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the most frequently recognized of the 31 epitopes that the vaccinated volunteers responded to.

HXB2 Location Nef (84–92)

**Author Location** Nef (84–92 BRU)

**Epitope** AVDLSHFLK

Subtype B, CRF02\_AG

Immunogen HIV-1 infection Species (MHC) human (A3, A11)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 0/9 CRF02\_AG-infected Ivorians, and 2/9 B-infected French subjects.
- One of the B-clade infected subjects that recognized this peptide carried the identical form, the other had one amino acid change: AIDLSHFLK. This variant form was in 3/5 B clade

infection sequences. 4/8 CRF01 infected individuals had a variant of this peptide.

HXB2 Location Nef (86–94)

**Author Location** Nef (84–92 LAI)

**Epitope** DLSHFLKEK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0301)

Keywords review

References McMichael & Walker 1994

· Review of HIV CTL epitopes.

HXB2 Location Nef (86–94)

**Author Location Nef** 

**Epitope** DLSHFLKEK **Subtype** A, B, D, F

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade

HIV component: p17 Gag, p24 Gag

**Species (MHC)** human, macaque (A\*0301)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee *et al.* [2002].

HXB2 Location Nef (86–94)

**Author Location** Nef (86–94)

Epitope DLSHFLKEK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A11, A\*0301)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot

granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- Two of nine patients responded to this peptide with GzB producing cells, while three of the patients responded with IFN-gamma producing cells. Only one patient had both GzB and IFN-gamma responses.

HXB2 Location Nef (86–94) Author Location Nef (86–94) Epitope DLSHFLKEK

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A3)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** Nef (86–94) **Author Location** Nef (86–94 HXB2)

Epitope DLSHFLKEK
Subtype B, CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A3, A11)

**Country** Viet Nam. **Assay type** HLA binding

**Keywords** subtype comparisons, computational epitope prediction, variant cross-recognition or cross-neutralization, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-All, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- The CRF01\_AE variant dlsFflkek had same HLA-binding score as the HXB2 epitope.

HXB2 Location Nef (86–100)

**Author Location** Nef (86–100 LAI)

Epitope DLSHFLKEKGGLEGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Robertson et al. 1993

Development of a retroviral vector (pNeoNef) to generate autologous targets.

HXB2 Location Nef (86–100)
Author Location Nef (86–100 LAI)
Epitope DLSHFLKEKGGLEGL
Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35) References Buseyne *et al.* 1993b

HXB2 Location Nef (86–100) Author Location Nef (86–100 LAI)

 ${\bf Epitope} \ \, {\tt DLSHFLKEKGGLEGL}$ 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35, Cw4)

References Buseyne et al. 1993a

- Vertical transmission of HIV ranges from 13% to 39%
- Primary assays showed that cytotoxic activity against at least one HIV protein was detected in 70% of infected children.
- Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.
- Patient EM13, who had a CTL response to three epitopes in Nef, was infected via blood transfusion after birth and went from CDC stage P2A to P2E during the study.

HXB2 Location Nef (87–102)

**Author Location** Nef

Epitope FSHFLKEKGGLEGLIY

Immunogen

Species (MHC) human

**Keywords** subtype comparisons **References** Jubier-Maurin *et al.* 1999

- 41 new HIV-1 strains describing envelope subtypes of HIV-1
   A-H were genetically characterized in the nef region 34 subtypes were classified in the same subtype in nef and env and 7 of the 41 strains were recombinants.
- This region was defined as a CTL epitope region that is conserved among HIV-1 M group subtypes.

HXB2 Location Nef (88-100)

**Author Location** Nef (103–116)

**Epitope SHFLKEKGGLEGL** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Guimaráes et al. 2002

 Nef sequences were obtained from Brazilians to study epitope diversity in this geographic region—most B subtype sequences are SHFLKEKGGLEGL, but sFflkekgglegl is found in most subtype C samples.

HXB2 Location Nef (90–97)

**Author Location** Nef (90–97)

**Epitope** FLKEKGGL

Epitope name BRU

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (A2, B8)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons **References** Inwoley *et al.* 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 1/9 CRF02\_AG-infected Ivorians, and 2/9 B-infected French subjects.

HXB2 Location Nef (90-97)

**Author Location Nef** 

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Keywords** dendritic cells

References Ostrowski et al. 2000

- The role of CD4+ T-cell help in expansion of virus-specific memory CTL was studied through co-culture ex vivo
- Optimal expansion of HIV-1-specific memory CTL depended on CD4+ T cell help in 9 of 10 patients – CD40 ligand trimer (CD40LT) could enhance CTL in the absence of CD4+ T cell help to a variable degree in most of patients.
- Those CTL that didn't respond to CD40LT could expand with IL2 present, and IL15 produced by dendritic cells also contributes.
- The T-helper epitope used for CD4+ T cell stimulation was the universal tetanus helper epitope TET830-843 (QYIKANSK-FIGITE)

HXB2 Location Nef (90–97)

**Author Location** 

Epitope FLKEKGGL

Epitope name Nef-FL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*08)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA B\*08, 1/3 (33%) recognized this epitope.

HXB2 Location Nef (90–97)

**Author Location** Nef (89–97 LAI)

Epitope FLKEKGGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*0801 epitope.

HXB2 Location Nef (90–97)

**Author Location** Nef (90–97)

Epitope FLKEKGGL

Epitope name FL8

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

```
      Donor MHC
      A*0201,
      A*2402,
      B*0801,
      B*5701,

      Cw*0602,
      Cw*0701;
      A*0101,
      A*0201,

      B*0801,
      B*5701,
      Cw*0602,
      Cw*0701;
      A2,

      A*2402,
      B*0801,
      B15,
      Cw7,
      Cw12;
      A*0101,

      A*0201,
      B*0801,
      B*5701,
      Cw*0602,

      Cw*0701
```

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFNγ, Tetramer binding, Chromium-release assay, Flow cytometric T-

cell cytokine assay

**Keywords** rate of progression, escape, TCR usage, characterizing CD8+ T cells

References Dong et al. 2004

- In 4 donors with delayed disease progression, the response
  to the FL8 Nef epitope was dominated by V-beta-13.2 TCR
  expressing CTLs with an unusually long CDR3 region. These
  CTLs were shown to be resistant to apoptosis and able to recognize escape variants of the FL8 Nef epitope. Thus, selection
  of these CTLs may be related to better clinical outcome.
- The Q5 variant flkeQggl was rapidly selected in a donor that responded to the FLKEKGGL epitope. The FLKEKGGL peptide and the variant flkeQggl HLA-B8 complexes bound to the Vbeta13.2 FLKEKGGL TCR with equal affinity, while the Vbeta6 FLKEKGGL TCR had reduced affinity for the FLKEKGGL form and did not recognize the Q5 variant. Other variants (T5, N5, and M5 as well as Q5) were recognized by Vbeta13.2 clones from all 4 donors. One clone from donor 046 that was not Vbeta13.2 could only recognize the index variant.

HXB2 Location Nef (90–97)

**Author Location** (C consensus)

**Epitope** FLKEKGGL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

HXB2 Location Nef (90–97)

**Author Location** Nef

Epitope FLKEKGGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0801)

**Donor MHC** A\*0101, B\*0801

Country United Kingdom.

Assav type CD8 T-cell Elispot - IFN $\gamma$ , HLA binding Keywords escape, acute/early infection References Milicic et al. 2005

- CTL responses and escape were analyzed in 4 homosexual couples where blood samples were available within weeks of estimated transmission. When the recipient had the same HLA type as the donor, CTL escape variants prevented a CTL response to those epitopes in the recipient. Even when the HLA alleles were different in the transmitting couple, a single escape mutation in one epitope can abolish CTL recognition of an overlapping epitope of distinct restriction in the recipient. In an early acute infection of the donor, the precise timing of transmission determines the viral variants transmitted.
- The recipient mounted an acute CTL response to this epitope, and the escape variant flkeQggl emerged soon after.

HXB2 Location Nef (90-97) **Author Location** (C consensus)

**Epitope** FLKEKGGL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0801) Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- FLKEKGGL is an optimal epitope.

HXB2 Location Nef (90–97)

**Author Location** Nef (89–97 LAI)

Epitope FLKEKGGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords review, escape

References Price et al. 1997

• CTL escape variants appeared over time in HLA-B8 HIV-1 + individual, providing evidence of immune escape.

- Most variants appear at position 5, an anchor residue.
- FLKE(E,N or Q)GGL showed reduced binding efficiency and recognition.
- Double mutants (FIKENGGL, FLEENGGL, and FLKGNGGL) completely escaped recognition.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study in the context of CTL escape to fixation.

HXB2 Location Nef (90-97)

**Author Location** Nef (90–97 IIIB)

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords HAART, ART, responses in children

References Spiegel et al. 1999

- Study examines the effect of highly active antiretroviral therapy (HAART) on HIV-1 plasma viral load, CTLp and CTLe frequencies in 8 infected children.
- CTLp (precursors) were measured by stimulating in culture and assaying using 51Cr release, against vaccina expressed IIIB Env, Gag, Pol, Nef.
- B7-FLKEKGGL tetramer complex was used for one of the children that was HLA-B7, and this infant showed a vigorous response (> 4% of CD8+ T cells) at 9 months of age.
- HIV-1 specific CTL responses initially increased in children with complete viral suppression, but then decreased, suggesting viral replication is needed to maintain CTL responses.

HXB2 Location Nef (90-97)

**Author Location Nef** 

Epitope FLKEKGGL

Immunogen vaccine

Vector/Type: vaccinia

Species (MHC) human (B8)

References Hanke et al. 1998a; Hanke et al. 1998b

• This epitope was shown to be processed and presented to appropriate CTL clones upon infection of human target cells with vaccinia virus Ankara (VVA) carrying 20 HIV-1 epitopes recognized by humans.

HXB2 Location Nef (90–97)

**Author Location** Nef (88–95)

**Epitope** FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Goulder et al. 1997g

- Natural variants for this epitope have been observed in several
- Substitutions Q5, N5, E5 that alter anchor position 5 are not well recognized.
- Substitution I2 binds well to B8 and is recognized.

HXB2 Location Nef (90–97)

**Author Location** Nef (90–97)

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Dyer et al. 1999

- CTL specific responses were measured over a 1.3 to 1.5 year period in members of the Sydney Blood Bank Cohort (SBBC) who had been infected with a natural attenuated strain of HIV-1 which was Nef-defective.
- Some of these patients had prolonged high levels of CTL effector and memory cells despite low viral load.

HXB2 Location Nef (90–97)

**Author Location** Nef (SF2)

Epitope FLKEKGGL

Epitope name FL8

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Goulder et al. 2001a

• This peptide elicited a weak CTL response during acute HIV-1 infection in patient PI004.

- Three CTL responses, to epitopes TSTLQEQIGW, ISPRTL-NAW, and KAFSPEVIPMF, were evident early after infection; CTL responses to SLYNTVATL, QASQEVKNW, EIYKRWII, and FLKEKGGL were detectable at 5 months post-infection and beyond.
- FL8 was recognized in an additional patient, AC29, in chronic infection.

**HXB2 Location** Nef (90–97) **Author Location** Nef (92–99)

Epitope FLKEKGGL

Epitope name FLK

Immunogen HIV-1 infection
Species (MHC) human (B8)
Keywords HAART, ART
References Oxenius *et al.* 2001a

- Characterization of specific CTL phenotype patterns in response to variation of the virus load in response to antiviral therapy in 3 patients with chronic HIV-1 infection.
- CTL activation in response to increasing viral load sequential, and co-segregated with apoptosis only during later stages of the response, suggesting antigen-specific cell-death is restricted to distinct CTL sub-populations.

HXB2 Location Nef (90–97)

Author Location Nef (92–99)

**Epitope** FLKEKGGL

Epitope name FLK

Immunogen HIV-1 infection Species (MHC) human (B8)

ies (MHC) human (B8)

Keywords HAART, ART, supervised treatment inter-

ruptions (STI), immunodominance, escape, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- Six of the 7/8 study subjects that were HLA B8 recognized this early dominant CTL epitope.
- Patient SC2 (HLA A1, B7/8, Cw0701/0702, DR4/53, DQ7) had CTL responsiveness against epitopes FLKEKGGL, GP-KVKQWPL, and GEIYKRWII peptides – FLKEKGGL tetramer staining steadily declined and at day 1340 the FLKEKGGL stained cells were no longer detected and the escape mutant FLKENGGI was found in 8/10 clones.
- Patient SC9 (HLA A1/2, B8/13, Cw0/0701, DR2/11, DQ6/7) had a CTL response against epitopes FLKEKGGL, ILKEPVHGV, SQRRQDILDLWIYHTQGYFPDWQNY, and GEIYKRWII peptides and all responses declined during therapy initiated at day 390 but were restored when therapy become intermittent.
- Patient SC19(HLA A11/12, B8/44, Cw06/0701, DR3/7, DR52/53, DQ 2/8) had a CTL response to epitopes

FLKEKGGL, GEIYKRWII, ACQGVGGPGHK, AVDLSH-FLK, and FNCGGEFFY that declined during therapy initiated at day 197.

- Patient SC10(HLA A1/3, B8/35, DR1/8, DQ 4/5) had sustained therapy started during acute infection and maintained an immunodominant response to FLKEKGGL and a response to GEIYKRWII through day 1088.
- Patient SC12(HLA A1, B8/39, Cw0701/0702, DR2/3, DR51/52, DQ2/6) had sustained therapy started during acute infection and maintained an immunodominant response to FLKEKGGL throughout and minor responses to GEIYKR-WII, DCKTILKAL, GGKKKYKLK – GEIYKRWII and GGKKKYKLK responses were stimulated by a brief period off therapy.
- Patient SC11(HLA A1, B8, Cw0201, DR3/11, DR52, DQ2/7) started therapy early, remained on therapy for 40 days, then reinitiated HAART at day 640 had a CTL response to FLKEKGGL, GPKVKQWPL, and GEIYKRWII throughout and received a benefit from the early limited course therapy.

HXB2 Location Nef (90–97)

**Author Location** Nef

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Kostense et al. 2001

- HLA tetramers to six epitopes were used to study HLA-A2, B8 and B57 CTL in 54 patients HIV-specific tetramer positive cells were inversely correlated with viral load in patients with high CD4, but in patients with CD4 T-cells below 400 high tetramer frequencies were found despite high viral load.
- Most patients have high levels of HIV-specific T-cell expansions, but many of these cells aren't functional.
- In 15 of the patients, the proportion of IFN gamma producing tetramer cells correlated with AIDS-free survival.
- Stimulation with HLA-B8 p24 and Nef epitopes significantly increased Nef-specific T-cell numbers in 2 patients (748 and 1113)
- There were more functional IFN-gamma producing Nefspecific T-cells within the T-cell population than there were active p24 Gag-specific T-cells.
- No correlation between elevated numbers of Nef-specific CTL cells and plasma viral load was observed.

HXB2 Location Nef (90–97)

**Author Location** Nef (88–95)

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

HXB2 Location Nef (90–97)

Author Location Nef (88–95 SF2)

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B8+ individuals that had a CTL response to this epitope broken down by group: 3/3 group 1, 1/3 group 2, and 1/2 group 3.

**HXB2 Location** Nef (90–97) **Author Location** Nef (89–97)

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Appay et al. 2000

- Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virusspecific CD8+ T cells produced IFN- $\gamma$  and MIP-1 $\beta$  with a distinct subset that failed to produce TNF- $\alpha$

**HXB2 Location** Nef (90–97) **Author Location** Nef (90–97)

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

References Day et al. 2001

- B8-restricted CTL accounted for about 1/3 of the total CTL response in one individual.
- The response to FLKEKGGL was the second highest response in magnitude compared to all the HLA class I A- and Brestricted epitopes tested in this individual.

HXB2 Location Nef (90–97)

**Author Location** Nef

Epitope FLKEKGGL

Immunogen HIV-1 infection Species (MHC) human (B8)

References Goulder et al. 2000b

- Tetramer assays were compared with three functional assays in 42 people with chronic HIV infection: ELISPOT, intracellular cytokine staining, and precursor frequency (limiting dilution assay [LDA])
- HIV-specific tetramer staining CTLs appeared to be active, and inert CTL were not found to play a significant role in chronic pediatric or adult HIV infection.

HXB2 Location Nef (90–97)

Author Location Nef (90–97 BRU)

**Epitope** FLKEKGGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Keywords** binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- FLKEKGGL was recognized in 12/14 (86%) of individuals with HLA B8, and it was a high affinity HLA binder.

HXB2 Location Nef (90–97)

**Author Location Nef** 

Epitope FLKEKGGL

Epitope name FLK

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI)

References Oxenius et al. 2002b

- Using previously defined epitopes Oxenius et al. [2000, 2001a] in an IFNgamma Elispot assay, 13 chronically HIV-1 infected patients were studied over a period including therapy with standard treatment interruptions (STI).
- STIs induced increased recognition of CTL epitopes, but there was no correlation between CTL responses with viral rebound rates, plateau viral loads, or clearance rates.

HXB2 Location Nef (90–97)

**Author Location Nef** 

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A2, A11, B8, B60, Bw6

Keywords HAART, ART

References Appay et al. 2002

- Four HIV patients with prolonged clinically successful antiviral therapy but with ongoing evidence of replication and Nef mRNA transcription, showed specific T-cell responses by Elispot and Tetramer staining, maintained for 2-4 years after initiation of HAART.
- Nef epitope recognition was detected in all 4 subjects, gp120,
   Pol and Gag-specific in 1 or 2 subjects two patients recognized FLKEKGGL.
- The HIV-specific CD8+ T-cells had an intermediate maturation phenotype characterized by low levels of perforin and high levels of CD27 expression.

HXB2 Location Nef (90–97)

**Author Location** Nef

Epitope FLKEKGGL Immunogen HIV-1 infection Species (MHC) human (B8)

**Donor MHC** A1, A3, B8, B65, Bw6

**Keywords** HAART, ART **References** Appay *et al.* 2002

- Four HIV patients with prolonged clinically successful antiviral therapy but with ongoing evidence of replication and Nef mRNA transcription, showed specific T-cell responses by Elispot and Tetramer staining, maintained for 2-4 years after initiation of HAART.
- Nef epitope recognition was detected in all 4 subjects, gp120, Pol and Gag-specific in 1 or 2 subjects – two patients recognized FLKEKGGL.
- The HIV-specific CD8+ T-cells had an intermediate maturation phenotype characterized by low levels of perforin and high levels of CD27 expression.

HXB2 Location Nef (90-97)

**Author Location Nef** 

**Epitope** FLKEKGGL **Subtype** A, B, C, D

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (B8)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

HXB2 Location Nef (90–97)

Author Location Nef (90-97)

**Epitope** FLKEKGGL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B8)

**Donor MHC** A1, A3, B8, B62, Cw3, Cw7 **Assay type** CD8 T-cell Elispot - IFNγ

**Keywords** binding affinity, acute/early infection, early-expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

HXB2 Location Nef (90–97)

Author Location Nef (90–97 B consensus)

Epitope FLKEKGGL

Epitope name FL8
Subtype B
Immunogen vaccine

Vector/Type: adeno-associated virus (AAV)

HIV component: gp120

Species (MHC) human (B8)

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay

Keywords dynamics, immune evasion

References Brainard et al. 2004

 HIV-1 gp120 is shown to suppress the ability of antigenspecific CTLs to migrate or remain at sites of high viral replication by concentration-dependent chemotaxis and fugetaxis. Directional T-cell movement is shown to depend on the interaction of the V2 and V3 loops with the CXCR4 receptor. X4 HIV-1 gp120 causes the migration of T-cells, including HIV-1 specific CTL, away from infected target cells, another potential mechanism for immune evasion.

HXB2 Location Nef (90–97)

**Author Location** Nef (87–95)

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A03, A28, B07, B08

Assay type proliferation, Chromium-release assay, Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, memory cells, immune dys-

function

References Gamberg et al. 2004a

• HAART restores HIV specific immunity after advanced infection by increase of CD4+ and CD8+ T cell numbers after supression of viral replication. However, HIV specific CTLs emerged only with detectable viral replication breakthroughs and were short-lived while CD4+ T-cell responses remained compromised, suggesting failure of generating stable CD8+ memory T-cells in the absence of HIV-specific T-helper responses.

HXB2 Location Nef (90-97)

**Author Location Nef** 

Epitope FLKEKGGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) (B8)

Keywords binding affinity, review, escape, characteriz-

ing CD8+ T cells

References da Silva 2003

- Evidence of the evolutionary adaptation of HIV-1 to the specific neutralizing antibody response and CTL detection is reviewed. Both SIV and HIV epitopes are discussed, with a detailed summary of one patient's response and CTL escape in the FLKEKGGL epitope. The three C-terminal amino acids were left unchanged, and it may be due to high fitness costs as these are putatively invloved in CD4 down-regulation and formation of a hydrophobic pocket in Nef. The N terminal residue is involved in binding to protein tyrosine kinases.
- Immediately after infection the susceptible eptiope FLKEKGGL was found in 20/20 viral sequences. Six months later, it was only found in 4/44 sequences. The flkeNggL form was most common, 24/44 cases; it bound poorly to HLA B08 and was poorly recognized by CTL. Two minor variants were found 3/44 times, flkeEggl and flkeQggl; both bound poorly to B08, but the K->Q substitution was still well recognized. A variant flkDkggl was found in 4/44 sequences; it bound B08 moderately well, but was poorly recognized. 3 double mutants were found once each, and were not recognized by CTL: flkeNggL, flEeNggL, and flkGNggL.

HXB2 Location Nef (90–97)

**Author Location** Nef (89–97)

**Epitope** FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human (B8)

Assay type Cytokine production, proliferation, CD8 T-

cell Elispot - IFNy, Tetramer binding, Intracellular cytokine staining, Flow cytometric T-cell cytokine assay

Keywords HAART, ART, memory cells, characterizing CD8+ T cells

References Daniel et al. 2004

• CD4+ and CD8+ responses in chronically HIV-1 infected patients on HAART were weak with decreased polyclonality. Only 33% of patients had CD4+ T-cells that could proliferate, and only 22% had HIV-specific CD8+ T-cells T-cell responses, and those rare responses showed low perforin levels and persistent expression of CD27, indicating incomplete differentiation and loss of lytic function.

HXB2 Location Nef (90-97)

**Author Location Nef** 

Epitope FLKEKGGL

Epitope name FL8

Immunogen HIV-1 infection

Species (MHC) human (B8)

Country United States.

Assay type proliferation, CD8 T-cell Elispot - IFNγ, Tet-

ramer binding

Keywords immunodominance, acute/early infection,

characterizing CD8+ T cells, immune dys-

function

References Lichterfeld et al. 2004a

- HIV-1 specific CD8+ T-cells in acute and long-term nonprogressive HIV-1 infection show strong ex-vivo proliferative capacities which are rapidly lost in chronic HIV-1 infection. The loss of CD8+ T-cell function is closely linked with the loss of HIV-1 specific, IL2 secreting CD4+ T-cells. The function can be rescued in vitro and in vivo by restoring the specific CD4+ T-cell help.
- Full CD8+ T-cell responses to this epitope were dependent on co-stimulation with a CD4+ T cell dependent epitope from T-cells harvested during acute infection. The CD8+ T-cell response to this epitope was immunodominant in one study individual.

HXB2 Location Nef (90–97)

Author Location (B consensus)

Epitope FLKEKGGL

Epitope name FL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A02, A03, B08, B62, Cw7, Cw10; A11, A29,

B08, B44, Cw4, Cw7; A25, A32, B08, B14, Cw7, Cw8; A01, A03, B08, B14, Cw7, Cw8

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.
- 4/9 individuals recognized this epitope, presented by HLA-B8.

HXB2 Location Nef (90-97)

**Author Location** Nef

Epitope FLKEKGGL

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B8)

Country United Kingdom.

Assay type Tetramer binding, T-cell Elispot, Intracellular

cytokine staining

acterizing CD8+ T cells, immune dysfunction

References Papagno et al. 2004

· Acute HIV-1 infection induces massive activation of HIVspecific and non-HIV-specific CD8+ T-cells resulting in differentiation of these cells. High differentiation of CD8+ T-cells is correlated with disease progression. Differentiation is a natural process but it can be driven by elevated immune activation, such as in HIV infection.

HXB2 Location Nef (90-97)

**Author Location Nef** 

**Epitope** FLKEKGGL

Epitope name FL8

Immunogen HIV-1 infection Species (MHC) human (B8)

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 5, FLKEnGGL, was found not to correspond to the most polymorphic residue in the epitope.

HXB2 Location Nef (90–97)

**Author Location Nef** 

Epitope FLKEKGGL

Epitope name FL8

Immunogen

Species (MHC) (B8)

**Keywords** review, immunodominance, acute/early infection, early-expressed

proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

• This review discusses the importance of 3 factors that impact the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides listed as immunodominant in acute HIV-1 infection.

HXB2 Location Nef (90-97)

**Author Location** Nef (90–97 HXB2)

Epitope FLKEKGGL

Epitope name FL8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B8)

**Donor MHC** A\*0101, A\*0201, B\*0801, B\*50, Cw\*0602,

Cw\*0701

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, reversion, viral fit-

ness, optimal epitope, HIV-1

References Liu et al. 2006

- **Keywords** rate of progression, acute/early infection, char• T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.
  - An FLKEeGGL presumed escape variant was transmitted from a B8 positive donor to a B8 negative recipient. FLKEgGGL, FLKgeGGL variants as well as FLKEkGGL reversions were found in the recipient subject.

HXB2 Location Nef (90-97)

**Author Location** Nef (89–97)

Epitope FLKEKGGL

Immunogen HIV-1 infection

Species (MHC) human

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INFγ responses to other epitopes.
- 1/11 of the A2+ individuals that responded to SLYNTVATL reacted with seven other epitopes including this epitope previously described as presented by B8.

HXB2 Location Nef (90-100)

Author Location Nef (90-100 BRU)

**Epitope** FLKEKGGLEGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Keywords** binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- FLKEKGGLEGL was recognized in 8/12 (67%) of individuals with HLA A2. It was a low affinity HLA A2 binder.

HXB2 Location Nef (90-104)

Author Location Nef (90-105 HXB2)

Epitope FLKEKGGLEGLIHSQ

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), immunodominance, early treatment

## References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%).
   p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- Responses to this peptide were detected in 16% of the study subjects, and it was one of the 25 most frequently recognized peptides.

HXB2 Location Nef (92-100)

**Author Location (LAI)** 

Epitope KEKGGLEGL

Subtype B

Immunogen

Species (MHC) human (B\*4001)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*4001,B60 epitope.

HXB2 Location Nef (92-100)

**Author Location** Nef

Epitope KEKGGLEGL

Epitope name KL9

Immunogen

**Species (MHC)** (B\*4001)

**Keywords** review, immunodominance, escape, acute/early infection, early-expressed

proteins, kinetics, reversion, viral fitness

References Lichterfeld et al. 2005

• This review discusses the importance of 3 factors that impact the selection of immunodominant epitopes in acute HIV infection: i) the kinetics of viral protein expression, ii) the HLA class I background of the infected individual, and iii) the autologous sequence of the infecting virus. This is 1 of 12 peptides listed as immunodominant in acute HIV-1 infection.

HXB2 Location Nef (92–100)

**Author Location** 

Epitope KEKGGLEGL

Epitope name Nef-KL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*4002)

**Donor MHC** A\*0201, A\*3201, B\*4002, B\*5301,

Cw\*0202, Cw\*0401

**Keywords** HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release
- Patient 01RCH59 was Hispanic, not on HAART, and had a viral load of 5100 and CD4 count of 349 – she also recognized TERQANFL, p2p7p1p6(64-70), HLA-B\*4002 and AEWDRVHPV, p24(78-86), HLA-B\*4002.
- Among HIV+ individuals who carried HLA B40, 3/5 (60%) recognized this epitope.

HXB2 Location Nef (92–100)

**Author Location** Nef (92–100)

Epitope KEKGGLEGL

Immunogen

Species (MHC) human (B\*4002)

Keywords optimal epitope

**References** Frahm et al. 2007

HXB2 Location Nef (92–100)

**Author Location** Nef (90–98 SF2)

Epitope KEKGGLEGL

Immunogen HIV-1 infection

Species (MHC) human (B60)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B60+ individuals that had a CTL response to this epitope broken down by group: 2/2 group 1, 1/1 group 2, and 0/0 group 3.

HXB2 Location Nef (92–100)

**Author Location** Nef (SF2)

Epitope KEKGGLEGL

Immunogen HIV-1 infection

Species (MHC) human (B60)

**References** Altfeld *et al.* 2000

This epitope was the dominant B60 (encoded by B\*4001) response in 6/8 HLA-B60 individuals, and recognized in all eight.

- This epitope was also recognized two expressing HLA-B61 individuals (B61 is usually encoded by B\*4002, but this study did not distinguish between B\*4002, B\*4003, B\*4004, B\*4006, and B\*4008)
- ELISPOT was a rapid an effective method that was used to define five novel B60 epitopes.
- HLA-B60 is present in 10-20% of the Caucasoid population and B60/B61 are very common in Asian populations.

HXB2 Location Nef (92-100)

**Author Location Nef** 

Epitope KEKGGLEGL Immunogen HIV-1 infection Species (MHC) human (B60) Keywords epitope processing References Cao et al. 2002

- KM is a B60 restricted CTL clone that recognizes KEKG-GLEGL.
- CTL could be activated by a fusion protein of an HIV protein and anthrax lethal factor (LFn-HIV) that promotes antigen presenting cell uptake of exogenous protein and allows processing through the MHC class I pathway. This strategy for CTL detection could allow antigen presentation without generation of cells by the standard methods of using live viral vectors carrying a protein, or by loading the cells with peptides and by-passing processing.

HXB2 Location Nef (92-100)

Author Location Nef (92–100 NL-43)

**Epitope** KEKGGLEGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B60)

**Keywords** class I down-regulation by Nef, escape

References Ali et al. 2003

- NL43 was passaged in the presence of Nef KEKGGLEGLspecific CTL clones STD11 and KM3, giving rise to rapid selection of escape mutations, including E93G, E93K, K94N+G99R, G95R+G99R, E98K, E98D, G99R, G99E, L100P, and L100I; insertions, deletions, frameshifts and an early stop codon. 34/36 (94%) of sequences carried mutations in the epitope by seven days, 36/36 (100%) by 14 days.
- NL43 was also passaged in the presence of a Nef TOGYFPDWONY-specific CTL clone. 7/15 clones had a frameshifting or stop codon introduced by one week; F121T was also observed. The most common escape mutation for both Nef epitopes was an early stop codon at position 91.
- Several mutations selected by KEKGGLEGL-specific CTL were shown to impair the down regulation of class I MHC by Nef, in particular E93G, E93K, and a truncation mutation at position 51.
- Nef deletion mutants increased 100-fold NL-43 susceptibility to inhibition by CTL specific for epitopes in other proteins, the A2 epitopes ILKEPVHGV in RT and SLYNTVATL in p17

HXB2 Location Nef (92-100)

**Author Location** Nef

Epitope KEKGGLEGL

Epitope name KL9

Immunogen HIV-1 infection

Species (MHC) human (B60)

**Donor MHC** A2, A24, B38, B60, Cw2, Cw12

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** supervised treatment interruptions (STI), early treatment

References Montefiori et al. 2003

• HIV-1 + patient AC10 underwent early HAART treatment, which was discontinued 1.5 years later. At this timepoint potent NAb responses against autologous virus were detected. Treatment interruption initially induced weak CD8+ responses directed against 5 epitopes. By days 873d and 923d the CTL response had broadened to target 22 epitopes; of these six were fully characterized. Eventually the virus escaped the NAb response, but escape was not accompanied by a rise in viral load, and the authors suggest the virus was contained by the CTL response.

HXB2 Location Nef (92–100)

**Author Location** Nef (92–100 NL43)

Epitope KEKGGLEGL

Epitope name KL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B60)

Assay type Chromium-release assay, CTL suppression of replication

Keywords escape

References Yang et al. 2003a

- · Virus was cultured in the presence of CTL lines specific for 5 different epitopes (SLYNTVATL, ILKEPVHGV, IEIKDTKEAL, SEGATPQDL, and KEKGGLEGL) to study the emergence of escape mutations. Escape varied between clones for the same epitope, and between different epitopes. Gag and RT epitope escape, if it occurred at all, tended to be monoclonal and within the epitope, indicating strong fitness constraints, while the Nef epitope escape was rapid, polyconal, and sometimes the result of upstream frameshifts.
- Two cloned CTL lines recongized KEKGGEGL, STD11 and KM3. Highly resistant clones emerged after a single round of passage with both CTL clones, and multiple substitutions accrued including frameshits and stop codons, reflecting the dispensability of Nef in viral culture.
- The following epitope variants were observed after passaging with clone STD11 for one week: kekggegI, kKkggegI, and 12/20 frameshifts and 1 early stop. By two weeks, a more complex polyclonal mixture was observed including: kekggegI, kKkggegl kekggegP, kekggeEl, kekggeRl, kekRgeRl, keNggeRl, and 11/22 framshifts.

HXB2 Location Nef (92–100)

Author Location Nef (92-100)

**Epitope** KEKGGLEGL

Epitope name Nef KL9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B60)

Assay type Chromium-release assay, Flow cytometric Tcell cytokine assay

CD8+ T cells

References Yang et al. 2003b

- Killing efficiency of CTLs and their ability to suppress viral replication is shown to depend more on epitope specificity than to antigenic avidity. Different clones recognizing the same epitope had similar killing efficiency despite their variation in avidity. Nef specific CTL clones tended to be most inhibitory, followed by Gag, then by RT specific clones, regardless of avidity.
- 2/14 CTL T-cell clones tested were specific for Nef KL9. Under conditions of excess peptide (100ug/ml), there was no difference in their lytic potential; all possessed similar effector capacity. Avidity was measured as the sensitizing dose of peptide required for 50% of maximal killing (SD50), which varied from 20 pg/ml to 100 ng/ml, over four orders of magnitude for all 14 epitopes. The SD50 range for Nef KL9 was 20-30 pg/ml, both high avidity. These clones were among the most efficient at inhibiting viral replication in the set tested, but because of the general lack of correlation between avidity and viral inhibition efficiency in this study, the authors attribute other reasons to Nefs ability to inhibit viral replication that pertain to presentation like kinetics and expression levels.

HXB2 Location Nef (92-100)

Author Location (B consensus)

Epitope KEKGGLEGL

Epitope name KL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B60) **Donor MHC** A03, B14, B60, Cw3, Cw7

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3 activation in dying target cells, it was shown that the subset of HIV-1-specific CD8+ T cells secreting both IFN-gamma and TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1specific CD8+ T-cell maturation phenotypes and intracellular perforin expression was found.
- 1/9 individuals recognized this epitope; the authors write that it is presented by HLA-B40 in their Table 1, but the subject that recognizes it, AC05, is HLA-B60, so we assume they meant B60.

HXB2 Location Nef (92-100)

**Author Location** Nef (92–100)

Epitope KEKGGLEGL

Immunogen HIV-1 infection

Species (MHC) human (B60, B61)

Keywords immunodominance

References Day et al. 2001

• No immunodominant responses were detected to five B61restricted epitopes tested.

**Keywords** binding affinity, TCR usage, characterizing • All five B60-restricted epitopes were reactive in another subject, and the B60-restricted responses together contributed over onethird of the total CTL response.

HXB2 Location Nef (92-105)

**Author Location Nef** 

Epitope KEKGGLEGLVYSQK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Novel unmapped epitope. There was a V->A change KEKG-GLEGLaYSQK over time in an individual that reacted with this peptide.

HXB2 Location Nef (92-112)

**Author Location** Nef (SF2)

Epitope KEKGGLEGLIHSQRRQDILDL

**Immunogen** HIV-1 infection

Species (MHC) human

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.
- The response to the peptide was CD8 dependent, but the HLA presenting molecule and optimal epitope were not determined.

HXB2 Location Nef (92-112)

**Author Location** Nef (SF2)

Epitope KEKGGLEGLIHSQRRQDILDL

Immunogen HIV-1 infection

Species (MHC) human

References Altfeld et al. 2000

- This epitope was mapped by ELISPOT in a study identifying new HLA-B60 epitopes, and was one of the epitopes presented by another HLA molecule in an HLA-B60 individual.
- The response to the peptide was CD8 dependent, but the HLA presenting molecule and optimal epitope were not determined.

HXB2 Location Nef (93–106)

**Author Location** Nef (93–106 BRU)

Epitope EKGGLEGLIHSQRR

Immunogen HIV-1 infection

Species (MHC) human (A1, B8) References Hadida et al. 1992

• HIV-1 specific CTLs detected in lymphoid organs of HIV-1 infected patients.

HXB2 Location Nef (97-111)

**Author Location** Nef (97–111)

Epitope LEGLIYSKKRQEILD

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** Nef (102–110)

**Author Location** Nef (102–110)

**Epitope** HSQRRQDIL

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (A\*24, B\*37, Cw\*0602, Cw\*0401)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope

prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** Nef (102–115)

Author Location Nef (102-115 LAI)

Epitope HSQRRQDILDLWIY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords review, escape

References Goulder et al. 1997e; Goulder et al. 1997a

- HLA identical sibling hemophiliac brothers were both infected with the same batch of factor VIII.
- They were tested 6-8 years after infection; one had a strong response to this peptide, the other did not.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study.

**HXB2 Location** Nef (102–115)

**Author Location** Nef (100–113)

Epitope HSQRRQDILDLWIY

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/7 patients recognized this epitope.

**HXB2 Location** Nef (102–121)

**Author Location** Nef (101–120 SF2)

Epitope HSQRRQDILDLQIYHTQGYF

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- Two of these 11 had CTL response to this peptide.
- The responding subjects were HLA-A2, A3, B8, B62 and HLA-A2, B21.

**HXB2 Location** Nef (103–117)

**Author Location** Nef

Epitope SKKRQEILDLWVYHT

Subtype A, D

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A\*0202, A\*3002, B\*5703, B\*5802; A\*6801,

A\*7401, B\*0702, B\*3501

Country Uganda.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, variant crossrecognition or cross-neutralization

References Barugahare et al. 2005

- T-cell immune responses were examined in HIV-1 infected Ugandans. Similar levels of cross-clade responses for Gag, Env and Nef were observed. Higher frequencies of responses in conserved regions were found for Gag within a single clade, while areas with higher viral sequence variation had fewer detectable responses. Differential interclade sequence homologies between the Gag regions did not affect the respective level of cross-clade recognition.
- The sequence contains a previously-defined epitope (KRQEILDLWVY) of unknown HLA restriction. The viral sequence from the subjects that recognized the peptide was skkrqKildlwvyNt.

**HXB2 Location** Nef (103–127)

Author Location Nef (103–127 PV22)

Epitope SQRRQDILDLWIYHTQGYFPDWQNY

Immunogen HIV-1 infection

**Species (MHC)** human (B13) **References** Jassoy *et al.* 1993

• HIV-1 specific CTLs release  $\gamma$ -IFN, and  $\alpha$ - and  $\beta$ -TNF.

HXB2 Location Nef (103–127) Author Location Nef (103–127)

Epitope SQRRQDILDLWIYHTQGYFPDWQNY

Epitope name SQR

Immunogen HIV-1 infection Species (MHC) human (B13)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- The only study subject out of eight that was HLA B13+ recognized this epitope.
- Patient SC9 (HLA A1/2, B8/13, Cw0/0701, DR2/11, DQ6/7) had a CTL response against epitopes FLKEKGGL, ILKEPVHGV, SQRRQDILDLWIYHTQGYFPDWQNY, and GEIYKRWII peptides and all responses declined during therapy initiated at day 390 but were restored when therapy become intermittent.

HXB2 Location Nef (104-112)

**Author Location** Nef (104–112)

Epitope QRRQDILDL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*37, Cw\*0602)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, computational epitope

prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** Nef (105–114)

Author Location Nef (105–114 LAI)

Epitope RRQDILDLWI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*2705)

Keywords rate of progression

References Goulder et al. 1997c

- Defined as optimal epitope from within reactive peptide HSQR-RQDILDLWIYHTQGYF [Nef(102-121 LAI)]
- HLA-B\*2705 is associated with slow HIV disease progression.

• The HLA-B\*2705 binding motif includes R at position 2, and L in the C-term position.

**HXB2 Location** Nef (105–114)

**Author Location** Nef (105–114 LAI)

Epitope RRQDILDLWI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*2705)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*2705 epitope.

**HXB2 Location** Nef (105–114)

**Author Location** Nef (105–114 SF2)

Epitope RRQDILDLWI

Immunogen HIV-1 infection

Species (MHC) human (B27)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B27+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 0/0 group 2, and 1/1 group 3.

**HXB2 Location** Nef (105–114)

**Author Location** Nef (105–114)

Epitope RRQDILDLWI

Immunogen HIV-1 infection

Species (MHC) human (B27)

**References** Day et al. 2001

 B27-restricted CTL response was strongest to this epitope in one individual.

**HXB2 Location** Nef (105–114)

**Author Location** 

Epitope RRQDILDLWI

**Epitope name** Nef-RI10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B27)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA B27, 1/2 (50%) recognized this epitope.

**HXB2 Location** Nef (105–115)

**Author Location** 

Epitope RRQDILDLWVY

Epitope name RY11 Immunogen

Species (MHC) human (B18)

Keywords optimal epitope References Frahm et al. 2007

• C. Brander notes this is a B18 epitope.

**HXB2 Location** Nef (105–115)

**Author Location** (C consensus)

Epitope KRQDILDLWIY

Subtype C

Immunogen HIV-1 infection Species (MHC) human (Cw\*0701)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the R2 residue of KRQDILDLWIY are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** Nef (105–115)

Author Location (C consensus)

Epitope KRQEILDLWVY

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (Cw\*0701, Cw\*0702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords cross-presentation by different HLA, characterizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** Nef (105–115)

**Author Location** (C consensus)

Epitope KRQDILDLWIY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- · Optimal epitope.

**HXB2 Location** Nef (105–115)

**Author Location** Nef (105–115)

Epitope RRQDILDLWIY

Epitope name Cw7-RY11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw7)

Donor MHC A3, B7, Cw7

Keywords dynamics, supervised treatment interruptions

(STI), acute/early infection

References Yu et al. 2002a

• AC-06 was homozygous at all three class I alleles (A3, B7, Cw7), and was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 response to RRQDILDLWIY restricted by HLA-Cw7.

**HXB2 Location** Nef (105–115)

Author Location Nef (105–115)

Epitope KRQEILDLWVY

Immunogen

Species (MHC) human (Cw7)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** Nef (105–115)

Author Location Nef (105–115)

Epitope RRQDILDLWIY

Immunogen

Species (MHC) human (Cw7)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** Nef (105–115)

**Author Location** Nef (C consensus)

Epitope KKQEILDLWVY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (Cw7)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape

References Kiepiela et al. 2004

• HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.

 People who carried Cw07 often carried a variant of this epitope, while the susceptible form of the epitope was highly conserved among those who did not.

HXB2 Location Nef (105–119)

Author Location Nef (105–119 HXB2)

Epitope RRQDILDLWIYHTQG

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

**Keywords** supervised treatment interruptions (STI), immunodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 19% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** Nef (106–114)

Author Location Nef (106–114)

Epitope RQDILDLWI

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (A\*24, B\*37, Cw\*0602)

Country India.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

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HXB2 Location Nef (106–114)
Author Location Nef (106–114)
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Epitope RQDILDLWI

Epitope name RI9

Immunogen HIV-1 infection Species (MHC) human (B13)

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, escape, variant crossrecognition or cross-neutralization, optimal

epitope

References Harrer et al. 2005

- An HLA-B13-restricted optimal epitope was defined in Nef, RI9. The frequency of CTLs specific for this epitope in B13positive patients exceeded the number of CTLs against other epitopes, indicating that this is a dominant epitope in B13positive subjects. Three B13-positive patients who had an immunodominant response to this epitope were good controllers of their infection, with low viral loads over long periods.
  - In B13-positive patients with a previous diagnosis of AIDS, an RrDILDLWI escape variant was found.
- This is a well conserved epitope but natural variants were tested. Peptide titation experiments indicate a V9I RQDILDLWv variant and RQDILDLWI are equally well recognized. Other natural substitutions are less well recognized: RkDILDLWI, RQeILDLWI, aQDILDLWI, RQaILDLWI, RrDILDLWv.

**HXB2 Location** Nef (106–114)

**Author Location** 

Epitope RQDILDLWV

Epitope name RV9

Immunogen

Species (MHC) human (B13)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes this is a B13 epitope.

**HXB2 Location** Nef (106–114)

Author Location Nef (106–114)

Epitope RQDILDLWI

Immunogen

**Species (MHC)** human (B13)

Keywords optimal epitope

**References** Frahm *et al.* 2007

• C. Brander notes that this is an B13 epitope.

**HXB2 Location** Nef (106–115)

**Author Location** (LAI)

Epitope RQDILDLWIY

Subtype B

Immunogen

**Species (MHC)** (B\*0702)

Keywords optimal epitope

References Frahm et al. 2007; Goulder 1999

**HXB2 Location** Nef (106–115)

**Author Location** Nef

Epitope RQDILDLWVY

Epitope name RY10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw7)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 9, RQDILDLWiY, was found in the most polymorphic residue in the epitope.

**HXB2 Location** Nef (107–115)

**Author Location** (C consensus)

Epitope QDILDLWIY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*18)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression **References** Kiepiela *et al.* 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in the D2 residue of QDILDLWIY are associated with the presence of the HLA presenting molecule in the host.
- QDILDLWIY not optimized.

**HXB2 Location** Nef (107–115)

Author Location Nef (107–115)

Epitope QDILDLWIY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*37, Cw\*0602)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

HXB2 Location Nef (108–115)

**Author Location** 

Epitope DILDLWIY

**Epitope name** Nef-DY8

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (Cw\*0701)

**Donor MHC** A\*3303 A\*2601 B\*5801 B\*8201 Cw\*0302

Cw\*0701

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described; 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Ct. release
- Subject 03RCH40 was African American, had a viral load of 2500, CD4 count of 372, was not on HAART, and also recognized the epitope ETKLGKAGY, RT(449-457), A\*2601.
- Among HIV+ individuals who carried HLA Cw07, 2/18 (11%) recognized this epitope.

**HXB2 Location** Nef (108–115)

**Author Location** Nef (108–115)

Epitope DILDLWIY

Immunogen HIV-1 infection

Species (MHC) human (Cw7)

**Donor MHC** A1, A1, B8, B14, Cw7, Cw8

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** binding affinity, acute/early infection, early-expressed proteins

References Cao et al. 2003

- CTL epitope responses were mapped in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized tended not to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently recognized than A1, A2, A30, and A44.

**HXB2 Location** Nef (109–117)

Author Location Nef (109–117)

Epitope ILDLWIYHT

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (A\*03)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** Nef (112–126)

**Author Location** Nef (112–126)

Epitope LWVYHTQGYFPDWQN

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Novitsky et al. 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 28 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** Nef (112–133)

**Author Location** Nef (111–132)

Epitope LWIYHTQGYFPDWQNYTPGPGV

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

• HIV-specific CTL lines developed by *ex vivo* stimulation with peptide.

**HXB2 Location** Nef (112–133)

**Author Location** Nef (111–132 SF2)

Epitope LWIYHTQGYFPDWQNYTPGPGV

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- Four of these 11 had CTL response to this peptide.
- The responding subjects were HLA-A2, B21; HLA-A1, A3, B7, B15; HLA-A2, A26, B7, B38.

**HXB2 Location** Nef (112–133)

**Author Location** Nef (111–132 SF2)

Epitope LWIYHTQGYFPDWQNYTPGPGV

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

 CTL expanded ex vivo were later infused into HIV-1 infected patients.

HXB2 Location Nef (113–121)

**Author Location** Nef (111–119)

Epitope WIYHTQGYF

Immunogen HIV-1 infection

Species (MHC) human (A1)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 3/13 patients recognized this epitope.

HXB2 Location Nef (113–121)

**Author Location** Nef (113–121)

Epitope WIYHTQGYF

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*35, Cw\*0602, Cw\*0401)

Country India.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

prediction, illimunodo

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** Nef (113–125)

Author Location Nef (113–125 BRU)

Epitope WIYHTQGYFPDWQ

Immunogen HIV-1 infection

Species (MHC) human (B17)

References Culmann et al. 1989

• Nef CTL clones from HIV+ donors.

HXB2 Location Nef (113–127)

**Author Location** Nef (128–142)

Epitope WIYHTQGYFDPWQNY

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Guimaráes et al. 2002

 Nef sequences were obtained from Brazilians to study epitope diversity in this geographic region – WIYHTQGYFDPWQNY displayed an (H) to (N) substitution in Brazilian Nef-gene subtype C samples, and this substitution is often found in other subtypes tested.

HXB2 Location Nef (113-128)

Author Location Nef (113–128 BRU)

Epitope WIYHTQGYFPDWQNYT

Immunogen HIV-1 infection

Species (MHC) human (A1)

References Hadida et al. 1992

HIV-1 specific CTLs detected in lymphoid organs of HIV-1 infected patients.

HXB2 Location Nef (113–128)

Author Location Nef (113–128 LAI)

Epitope WIYHTQGYFPDWQNYT

Epitope name N2

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A1)

Keywords HAART, ART

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+ PBL but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

HXB2 Location Nef (114–127)

**Author Location** Nef

Epitope VYHTQGYFPDWQNY

Immunogen HIV-1 infection

Species (MHC) human

References Jubier-Maurin et al. 1999

**HXB2 Location** Nef (115–125)

Author Location Nef (115-125 BRU)

Epitope YHTQGYFPDWQ

Immunogen HIV-1 infection

Species (MHC) human (B17)

References Culmann et al. 1991

• Nef CTL clones from HIV+ donors.

**HXB2 Location** Nef (115–129)

614

**Author Location** Nef (115–129 HXB2)

 ${\bf Epitope} \ \ {\tt YHTQGYFPDWQNYTP}$ 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 22% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** Nef (116–124)

Author Location Nef (116–124)

Epitope HTQGYFPDW

Immunogen

Species (MHC) human (B\*57)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** Nef (116–124)

**Author Location** Nef (116–124)

Epitope HTQGYFPDW

Immunogen

Species (MHC) human (B57)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** Nef (116–124)

**Author Location** 

Epitope HTQGYFPDW

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** assay standardization/improvement, epitope

processing

References Draenert et al. 2004a

- 96% of optimally defined epitopes have one of only nine amino acids serving as the C-terminal anchor position. Seven amino acids are never found in this position and four are only present in 4% of cases. CD8 T-cell response to an epitope is shown to be best detected when the epitope is situated at the C-terminal end of a longer peptide, and authors suggest that Elispot reagents would be better designed if peptides ended on known C-terminal anchors.
- HTQGYFPDW is suggested to be the optimal epitope instead of HTQGYFPDWQ since Gln is not described as a C-terminal anchor residue in any of the other optimally defined epitopes. HTQGYFPDW was also found to be recognized at two times lower peptide concentrations than HTQGYFPDWQ.

**HXB2 Location** Nef (116–124)

Author Location Nef (116–124)

Epitope HTQGYFPDW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B57)

**Donor MHC** A\*3001, A\*66, B\*4201, B\*5802, Cw\*0602, Cw\*1701; A\*68, A\*66, B\*57, B\*5802,

Cw\*0602, Cw\*0701

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords responses in children, mother-to-infant trans-

mission, escape, acute/early infection

References Pillay et al. 2005

- HIV-specific CTLs in infants were shown to be able to select for viral escape variants early in life, despite a lack of escape with the same CTL specificity in the mother. Infant CTL responses may be compromised by transmission of escape variants that arose in the mother and also those that arose in the father, if the father was the source of the mother's infection.
- HTQGYFPDW is the C consensus form of the epitope; the autologous form in the mother was HTQGfFPDW, and this was transmitted to her infant. By 33 weeks a new dominant form of the epitope had emerged in the infant: nTQGfFPDW.

**HXB2 Location** Nef (116–124)

**Author Location Nef** 

Epitope HTQGYFPDW

Epitope name HW9

Immunogen HIV-1 infection

Species (MHC) human (B57, B\*5801)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release

Keywords responses in children, mother-to-infant trans-

mission, escape, characterizing CD8+ T cells

References Feeney et al. 2005

• Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and

that a developing immune system may exhibit greater plasticity in recognizing viral variants.

While 2 mothers carried the form HTQGYFPDW, their children carried the escape H1N variant nTQGYFPDW.

**HXB2 Location** Nef (116–124)

Author Location Nef (116–124 BRU)

Epitope HTQGYFPDW

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (B57, B58)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 0/9 CRF02\_AG-infected Ivorians, and 1/9 B-infected French subject
- HTQGYFPDW was invariant in 5 B clade infected individuals, including the one that recognized the epitope. It varied in 4/8 CRF02 Ivorian infections.

**HXB2 Location** Nef (116–125)

Author Location Nef (116-125 BRU)

Epitope HTQGYFPDWQ

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords subtype comparisons, optimal epitope

References Frahm et al. 2007

- C. Brander notes this is a B\*5701 epitope.
- Subtype of B57 not determined.

HXB2 Location Nef (116-125)

**Author Location** Nef (116–125)

Epitope HTQGYFPDWQ

Immunogen HIV-1 infection

Species (MHC) human (B57)

Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- One of the A2+ individuals was HLA A\*0201, A1, B57 and responded to four B57 epitopes and two others.

**HXB2 Location** Nef (116–125)

**Author Location** Nef (116–125 BRU)

**Epitope** HTQGYFPDWQ

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Culmann et al. 1991

Nef CTL clones from HIV+ donors, optimal peptide mapped.

HXB2 Location Nef (116-125)

Author Location Nef (116-125)

Epitope HTQGYFPDWQ

Epitope name HTQ

Immunogen HIV-1 infection Species (MHC) human (B57)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- None of the 8 study subjects recognized this epitope but none were HLA B57+

**HXB2 Location** Nef (116–125)

**Author Location** 

Epitope HTQGYFPDWQ

Epitope name Nef-HQ10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA B57, 0/5 (0%) recognized this epitope.

HXB2 Location Nef (116-125)

Author Location Nef (114-123)

Epitope HT0GYFPDW0

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 7 patients recognized this epitope.

**HXB2 Location** Nef (116–125)

**Author Location Nef** 

Epitope HTQGYFPDWQ

**Epitope name** HQ10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B57)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 1, nTQGYFPDWQ, was found in the most polymorphic residue in the epitope.

**HXB2 Location** Nef (116–125)

**Author Location Nef** 

Epitope HTQGYFPDWQ

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human (B57, B63)

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, cross-presentation by dif-

ferent HLA, optimal epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This epitope was recognized by 30% of B63-positive subjects and 35% of B57/58-positive subjects.

**HXB2 Location** Nef (117–127)

Author Location Nef (117–127 LAI)

Epitope TQGYFPDWQNY

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human (B\*1501)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1501 epitope.

HXB2 Location Nef (117-127)

Author Location Nef (117–127 NL-43) Epitope TQGYFPDWQNY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1501)

Keywords class I down-regulation by Nef, escape

References Ali et al. 2003

NL43 was passaged in the presence of Nef KEKGGLEGL-specific CTL clones STD11 and KM3, giving rise to rapid selection of escape mutations, including E93G, E93K, K94N+G99R, G95R+G99R, E98K, E98D, G99R, G99E, L100P, and L100I; insertions, deletions, frameshifts and an early stop codon. 34/36 (94%) of sequences carried mutations in the epitope by seven days, 36/36 (100%) by 14 days.

- NL43 was also passaged in the presence of a Nef Author Location Nef (117–147 LAI) TQGYFPDWQNY-specific CTL clone. 7/15 clones had a frameshifting or stop codon introduced by one week; F121T was also observed. The most common escape mutation for both Nef epitopes was an early stop codon at position 91.
- Several mutations selected by KEKGGLEGL-specific CTL were shown to impair the down regulation of class I MHC by Nef, in particular E93G, E93K, and a truncation mutation at position 51.
- Nef deletion mutants increased 100-fold NL-43 susceptibility to inhibition by CTL specific for epitopes in other proteins, the A2 epitopes ILKEPVHGV in RT and SLYNTVATL in p17 Gag.

**HXB2 Location** Nef (117–127) **Author Location** Nef (117–127 LAI) Epitope TQGYFPDWQNY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B62) References Culmann 1998

• Optimal peptide defined by titration.

HXB2 Location Nef (117–127) **Author Location** Nef (117–127) Epitope TQGYFPDWQNY Immunogen HIV-1 infection Species (MHC) human (B62) Keywords immunodominance

References Day et al. 2001

• No immunodominant responses were detected to four B62restricted epitopes tested.

HXB2 Location Nef (117–127) **Author Location** Nef (117–127) **Epitope** TQGYFPDWQNY Immunogen HIV-1 infection Species (MHC) human Keywords immunodominance

References Betts et al. 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INF $\gamma$  responses to other epitopes.
- 1/11 of the A2+ individuals was HLA A\*0205/A\*0208, A30, B27, B44 but responded to HLA Bw62 epitope TQGYF-PDWQNY, and one of the other individuals that was A2+, but otherwise of unknown HLA type, reacted with seven epitopes including this one.

**HXB2 Location** Nef (117–128) **Author Location** Nef (117–128 BRU) **Epitope** TQGYFPDWQNYT Immunogen HIV-1 infection Species (MHC) human (B17, B37) References Culmann et al. 1991

· Nef CTL clones from HIV+ donors.

HXB2 Location Nef (117–147)

Epitope TQGYFPDWQNYTPGPGVRYPLTFGWCYKLVP

Subtype B Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long amino acid peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees - 1/10 reacted to this Nef peptide.
- 9/12 tested mounted a CTL response to at least one of the six peptides; each of the six peptides elicited a CTL response in at least one individual.
- 10/12 tested had an IgG response to this peptide.

HXB2 Location Nef (118-127) **Author Location** Nef (118–127 LAI) Epitope QGYFPDWQNY Subtype B Immunogen

Species (MHC) human (B62)

Keywords review

References McMichael & Walker 1994

• Review of HIV CTL epitopes.

**HXB2 Location** Nef (119–127) **Author Location** Nef (119–127) **Epitope** GYFPDWONY Subtype B, CRF02 AG Immunogen HIV-1 infection Species (MHC) human (A24) Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ **Keywords** subtype comparisons References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 0/9 CRF02\_AG-infected Ivorians, and 1/9 B-infected French subjects. It was invariant among 5 French subjects, including the one that reacted with the epitope, and had single amino acid substitutions in 4/8 Ivorians.

**HXB2 Location** Nef (120–127) **Author Location** (C consensus)

Epitope YFPDWQNY

Subtype C

Immunogen HIV-1 infection Species (MHC) human (A\*29)

Country South Africa. Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- YFPDWQNY is an optimal epitope.

HXB2 Location Nef (120-127)

Author Location (C consensus)

Epitope YFPDWQNY

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (A\*3002, A\*29)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords cross-presentation by different HLA, charac-

terizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** Nef (120–127)

**Author Location** 

**Epitope** YFPDWQNY

Immunogen HIV-1 infection Species (MHC) human (A29)

Country United States.

Assav type CD8 T-cell Elispot - IFNγ

Keywords assay standardization/improvement, epitope

processing

References Draenert et al. 2004a

- 96% of optimally defined epitopes have one of only nine amino acids serving as the C-terminal anchor position. Seven amino acids are never found in this position and four are only present in 4% of cases. CD8 T-cell response to an epitope is shown to be best detected when the epitope is situated at the C-terminal end of a longer peptide, and authors suggest that Elispot reagents would be better designed if peptides ended on known C-terminal anchors.
- Instead of YFPDWQNYT, YFPDWQNY was found to be the optimal epitope in one patient.

**HXB2 Location** Nef (120–127)

**Author Location** 

**Epitope** YFPDWQNY Immunogen HIV-1 infection

Species (MHC) human (B57, B\*5801)

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assav

Keywords responses in children, mother-to-infant transmission, escape

References Feeney et al. 2005

- Escape mutations in TW10 and other B57-restricted epitopes were shown to arise early in infants following perinatal infection. Some escape variants were likely to have been transmitted vertically, from HLA-B57/5801 positive HIV-1 infected mothers, while others arose during infancy in cases where the children inherited the B57/5801 allele paternally. In contrast to adults, the majority of children showed a robust response to the escape variants, suggesting that infants are able to mount functional immune responses and drive immune escape and that a developing immune system may exhibit greater plasticity in recognizing viral variants.
- YFPDWQNY responses were somewhat more frequent in adults.

**HXB2 Location** Nef (120–128)

**Author Location** Nef (118–126 SF2)

Epitope YFPDWQNYT

Immunogen HIV-1 infection

Species (MHC) human (A1)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- · Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A1+ individuals that had a CTL response to this epitope broken down by group: 0/3 group 1, 0/2 group 2, and 1/2 group 3.

**HXB2 Location** Nef (120–128)

**Author Location** Nef (120–128)

Epitope YFPDWQNYT

Immunogen HIV-1 infection

Species (MHC) human (A29)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** Nef (120–128)

**Author Location** Nef (120–128 LAI)

Epitope YFPDWQNYT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*3701)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*3701 and B\*5701 epitope.

**HXB2 Location** Nef (120–128)

Author Location Nef (120-128 LAI)

Epitope YFPDWQNYT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5701)

Keywords optimal epitope

References Frahm et al. 2007

- C. Brander notes this is a B\*5701 epitope.
- Subtype of B57 not determined.

**HXB2 Location** Nef (120–128)

Author Location Nef (120-128 IIIB)

Epitope FFPDWKNYT

Immunogen HIV-1 infection

Species (MHC) human (B15)

Keywords responses in children, mother-to-infant trans-

mission, escape

References Wilson et al. 1999a

- This study describes maternal CTL responses in the context of mother-to-infant transmission.
- Detection of CTL escape mutants in the mother was associated with transmission, but the CTL-susceptible forms of the virus tended to be found in infected infants.
- · LFPDWKNYT is an escape mutant.

HXB2 Location Nef (120-128)

Author Location Nef (120-128 LAI)

Epitope YFPDWQNYT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B37, B57)

References Culmann 1998

 Nef CTL clones from HIV+ donors – optimum peptide mapped by titration.

**HXB2 Location** Nef (120–128)

**Author Location** Nef (120–128)

Epitope FFPDWKNYT

Immunogen HIV-1 infection

Species (MHC) human (B51)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 5 epitopes from this individual that varied over time where the internal mutations became fixed; a dramatic decrease in CTL activity against the wild type epitope was observed as the mutation arose. YfpdwQnyt and YfpdwHnyt variants found at 2 months postseroconversion (psc); YfpdwHnyt, YfpdwQSyt, YLpdwQSyt and YfpdwDnyt variants found 20 months psc; YfpdwDnyt and YfpdwQSyt variants found 47 months psc.

**HXB2 Location** Nef (120–128)

**Author Location** 

Epitope YFPDWQNYT

**Epitope name** Nef-YT9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Sabbaj et al. 2003

 Among HIV+ individuals who carried HLA B57, 1/5 (20%) recognized this epitope.

HXB2 Location Nef (120-128)

**Author Location Nef** 

Epitope YFPDWQDYT

**Epitope name** YT9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B57)

Donor MHC A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

 $\textbf{Keywords} \ \ \text{subtype comparisons, escape, characterizing}$ 

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- One escape mutation, at position 7, YFPDWQnYT, was found not to correspond to the most polymorphic residue in the epitope.

**HXB2 Location** Nef (120–128)

**Author Location Nef** 

Epitope YFPDWQNYT

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human (B57, B63)

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, cross-presentation by dif-

ferent HLA, optimal epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58.
- This epitope was recognized by 40% of B63-positive subjects and 27% of B57/58-positive subjects.

HXB2 Location Nef (120–128)

**Author Location** 

Epitope YFPDWQNYT

Immunogen

Species (MHC) human (Cw6)

Keywords optimal epitope
References Frahm *et al.* 2007

• C. Brander notes that this is an Cw6 epitope.

HXB2 Location Nef (120–128)
Author Location Nef (120–128)
Epitope YFPDWQNYT
Immunogen HIV-1 infection
Species (MHC) human

**Keywords** immunodominance **References** Betts *et al.* 2000

- Only 4/11 HLA-A2+ HIV+ individuals had CTL that reacted to SLYNTVATL, calling into question whether it is immunodominant.
- 95 optimally-defined peptides from this database were used to screen for INFγ responses to other epitopes.
- 1/11 of the A2+ individuals was HLA A\*0205/A\*0208, A30, B27, B44 but responded to HLA B37 epitope IYKRWIILGL, and one of the other individuals that was A2+, but otherwise of unknown HLA type, reacted with seven epitopes including this one.

**HXB2 Location** Nef (120–144) **Author Location** Nef (120–144 SF2)

Epitope YFPDWQNYTPGPGIRYPLTFGWCYK

Immunogen HIV-1 infection Species (MHC) human (A24) References Jassoy *et al.* 1992

· Epitope recognized by CTL clone derived from CSF.

HXB2 Location Nef (121–128) Author Location Nef (121–128) Epitope FPDWQNYT Subtype B Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (A1)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of 31 that the vaccinated volunteers responded to.

HXB2 Location Nef (121–128)

Author Location Nef (121–128 HXB2)

Epitope FPDWQNYT

Subtype B, CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A1)

**Country** Viet Nam. **Assay type** HLA binding

**Keywords** subtype comparisons, computational epitope prediction, variant cross-recognition or cross-neutralization, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-All, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- CRF01\_AE has 2 forms, FPDWQNYT, like the HXB2 form, and FPDWhNYT. Both are predicted to bind to A1.

HXB2 Location Nef (122–136) Author Location Nef (122–136)

Epitope PDWQNYTPGPGVRYP

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 8 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

**HXB2 Location** Nef (122–141) **Author Location** Nef (121–140 SF2)

Epitope PDWQNYTPGPGVRYPLTFGW

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- Three of these 11 had CTL response to this peptide.
- The responding subjects were HLA-A2, B21; HLA-A3, A24, B7, B38.

HXB2 Location Nef (123–137)

Author Location Nef (123–137 IIIB)

Epitope QWQNYTPGPGVRYPL

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** responses in children, mother-to-infant transmission, escape

**References** Wilson et al. 1996

- Epitope defined in the context of the Pediatric AIDS Foundation ARIEL Project, a mother-infant HIV transmission study.
- FFPDYTPGPGTRFPL and FFPDYKPGPGTRFPL, naturally occurring variants, were found in mother and are not recognized.

 LFPDYKPGPGTRFPL and FFPDYKPGPGTRFPL, naturally occurring variants, were found in infant and are not recognized.

HXB2 Location Nef (126–135) Author Location Nef (126–135 BRU) Epitope NYTPGPGVRY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A24)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- NYTPGPGVRY was recognized in 3/10 (30%) of individuals with HLA A24. It was a moderate affinity HLA-A24 binder.

**HXB2 Location** Nef (126–138)

Author Location Nef (126–138 BRU)

Epitope NYTPGPGVRYPLT

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Culmann et al. 1991

· Nef CTL clones from HIV+ donors.

**HXB2 Location** Nef (127–135)

**Author Location** 

Epitope YTPGPGIRY

Immunogen

Species (MHC) human (B57)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes that this is an B57 epitope.

**HXB2 Location** Nef (127–135)

**Author Location** Nef

Epitope YTPGPGIRY

Epitope name YY9

Subtype B, C

Immunogen HIV-1 infection Species (MHC) human (B57, B58)

**Donor MHC** A02, A33, B35, B57, Cw04, Cw07

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

 HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and B58

• This is a putative HLA-B63/57/58 epitope containing the B58 supertype binding motif. The optimal epitope was defined in a person carrying B57, and reactivity to the peptide was enriched in those with B57/B58, just a trend for B63.

**HXB2 Location** Nef (127–135)

**Author Location** 

Epitope YTPGPGIRY

Immunogen

Species (MHC) human (B63)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B63 epitope.

**HXB2 Location** Nef (127–141)

**Author Location** Nef (127–141)

Epitope YTPGPGVRYPLTFGW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

**References** Novitsky *et al.* 2002

- HLA, viral sequence, and Elispot data was obtained from 105 HIV-1 positive Botswanans; Elispot data was obtained from between 55 and 64 subjects for each HIV protein.
- Nef and p24 had the highest percentage of reactive peptides, and p24 had the highest magnitude of HIV-1 responses.
- This peptide was among the 8 most reactive C clade peptides from among over 350 tested spanning all HIV proteins.

HXB2 Location Nef (128–135)

Author Location Nef (128–135 LAI)

Epitope TPGPGVRY

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (B\*0702)

Keywords epitope processing

References Lucchiari-Hartz et al. 2000

- Five naturally processed MHC class I ligands were identified in Nef in the conserved immunogenic region Nef between 123-152.
- All five could be transported by TAP, and 4/5 had N-termini that were major cleavage points for the proteasome, only one had extended precursor fragments.
- Both TPGPGVRYPL and TPGPGVRY are naturally processed ligands that can be eluted from HLA-B7 molecules, both are recognized by the same CTL, and both peptides seem to be the direct product of a proteasomal digest.
- The peptide TPGPGVRY is present in a high copy number, TPGPGVRYPL at a more moderate level, possibly due to a major cleavage site between the Y and P.

**HXB2 Location** Nef (128–135)

**Author Location** Nef (128–135)

Epitope TPGPGVRY

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env. Gag. Nef Adjuvant: OS21

Species (MHC) human (B7 supertype)

Assay type proliferation, CD8 T-cell Elispot - IFNγ, Chromium-release assay

Keywords vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

• After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

**HXB2 Location** Nef (128–136)

**Author Location** 

Epitope TPGPGVRYP

Epitope name Nef-TP9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA B07, 4/9 (44%) recognized this epitope.

**HXB2 Location** Nef (128–137) Author Location Nef (128–137 LAI)

**Epitope** TPGPGVRYPL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

**Keywords** optimal epitope References Frahm et al. 2007

• C. Brander notes this is a B\*0702 epitope.

HXB2 Location Nef (128-137) Author Location Nef (128–137 LAI)

Epitope TPGPGVRYPL

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (B\*0702)

Keywords epitope processing

References Lucchiari-Hartz et al. 2000

- Five naturally processed MHC class I ligands were identified in Nef in the conserved immunogenic region Nef between 123-152.
- All five could be transported by TAP, and 4/5 had N-termini that were major cleavage points for the proteasome, only one had extended precursor fragments.
- Both TPGPGVRYPL and TPGPGVRY are naturally processed ligands that can be eluted from HLA-B7 molecules, both are recognized by the same CTL, and both peptides seem to be the direct product of a proteasomal digest.
- The peptide TPGPGVRY is present in a high copy number, TPGPGVRYPL at a more moderate level, possibly due to a major cleavage site between the Y and P.

**HXB2 Location** Nef (128–137)

Author Location (C consensus)

**Epitope** TPGPGVRYPL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*0702)

Country South Africa.

Assav type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- TPGPGVRYPL is an optimal epitope for B\*4201 and B\*0702.

**HXB2 Location** Nef (128–137)

Author Location Nef (127–136)

**Epitope** TPGPGVRYPL

Epitope name TL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*3501)

Donor MHC A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, Chromium-release assay

Keywords escape, acute/early infection

References Bansal et al. 2005

- · Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.
- Point mutation at position 2 (P to S, TsGPGVRYPL) was detected at a chronic infection time point, month 33. This escape variant had lower avidity.
- · The response to the peptide that carried this epitope was initially strong and diminished over time.

HXB2 Location Nef (128–137)

**Author Location** Nef (128–137 LAI)

**Epitope** TPGPGVRYPL

Subtype B

Immunogen

Species (MHC) human (B\*4201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*4201 epitope.

HXB2 Location Nef (128-137)

**Author Location** Nef (128–137)

Epitope TPGPGVRYPL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

**Donor MHC** A\*3001, A\*30, B\*1503, B\*4201, Cw\*0202,

Cw\*1701; A\*0301, A\*3001, B\*4201, **Author Location** Nef (128–137 BRU)

B\*5802, Cw\*0602, Cw\*1701

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection

References Pillay et al. 2005

- HIV-specific CTLs in infants were shown to be able to select for viral escape variants early in life, despite a lack of escape with the same CTL specificity in the mother. Infant CTL responses may be compromised by transmission of escape variants that arose in the mother and also those that arose in the father, if the father was the source of the mother's infection.
- Escape variants tSgpgvrypl and tQgpgvrypl were rapidly selected in an infant, by 26 weeks, but were not found in the infant's mother. These forms were demonstrated to be escape mutations by Elispot, and also had reduced binding to B\*4201 in a competitive inhibition assay.

HXB2 Location Nef (128-137)

**Author Location** (C consensus)

Epitope TPGPGVRYPL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*4201)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.

**HXB2 Location** Nef (128–137)

**Author Location** (C consensus)

Epitope TPGPGVRYPL

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*4201, B\*0702)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords cross-presentation by different HLA, charac-

terizing CD8+ T cells

References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1 were analyzed in African patients. Significantly more responses were shown to be HLA-B restricted. Viral load, CD4 count, and thus rate of disease progression were also associated with HLA-B alleles. In addition, the selection pressure imposed on HIV-1 by HLA-B alleles was shown to be substantially greater than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** Nef (128–137)

**Epitope** TPGPGVRYPL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

**Keywords** binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- TPGPGVRYPL was recognized in 8/16 (50%) of individuals with HLA B7, and 1/9 (11%) of individuals with HLA B35. It was a high affinity HLA binder.

HXB2 Location Nef (128-137)

**Author Location** 

**Epitope** TPGPGVRYPL

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords acute/early infection

References Wilson et al. 2000a

- · Three individuals with highly focused HIV-specific CTL responses were studied during acute infection using tetramers high frequencies of HIV-1-specific CD8+ T cells were found prior to seroconversion, and there was a close temporal relationship between the number of circulating HIV-specific T cells and viral load was also found.
- TPGPGVRYPL is an optimal epitope for B\*4201 and B\*0702. All three patients were B\*2705, with HLA alleles: A1, A30/31, B\*2705, B35; A1, A\*0301, B7, B\*2705; and A\*0201, A\*0301, B\*2705, B39.
  - ELISPOT was used to test a panel of CTL epitopes that had been defined earlier and were appropriate for the HLA haplotypes of the study subjects -3/3 subjects showed a dominant response to the B\*2705 epitope KRWIILGGLNK.
  - The subject with A\*0201 had a moderately strong response to SLYNTVATL.
  - Weak responses were observed to A\*301-RLRPGGKKK, A\*301-QVPLRPMTYK, and B7-TPGPGVRYPL in the subject who was HLA A1, A\*0301, B7, B\*2705.
  - · No acute response was detected to the following epitopes: A\*201-ILKEPVHGV, A\*301-KIRLRPGGK, A\*301-AIFQSSMTK, A\*301-TVYYGVPVWK, B35-EPIVGAETF, B35-HPDIVIYQY, B35-PPIPVGEIY, B35-NSSKVSQNY, B35-VPLRPMTY, B35-DPNPQEVVL.

**HXB2 Location** Nef (128–137)

**Author Location** Nef (128–137 LAI)

Epitope TPGPGVRYPL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Haas et al. 1996; Haas et al. 1997

- There was a high degree of variation in three CTL epitopes in Nef in four slow and non-progressors, and variant specific CTLs arose over time to eliminate variants, indicating immune selection.
- The epitope position was taken from Haas et al. [1997]

**HXB2 Location** Nef (128–137)

**Author Location Nef** 

Epitope TPGPGVRYPL

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B7)

**Keywords** subtype comparisons, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades such cross-reactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The D subtype consensus is identical to the B clade epitope.
- The A subtype consensus is TPGPGIRYPL.

**HXB2 Location** Nef (128–137)

**Author Location** Nef (subtype B)

Epitope TPGPGVRYPL

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B7)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found B clade epitopes are often cross-reactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among B and D clade viruses.
- The Clade A version of the epitope: TPGPGIRYPL.

HXB2 Location Nef (128–137)

**Author Location** Nef (128–137)

**Epitope** TPGPGVRYPL

Immunogen in vitro stimulation or selection

Species (MHC) human (B7)

Keywords immunodominance, dendritic cells, Th1

References Wilson et al. 1999b

- Dendritic cells are the most potent for priming T cell responses
   DCs can stimulate autologous CTL responses from T cells cultured from HIV negative donors.
- Th1-biasing cytokines IL-12 or IFN alpha enhance CTL responses in vitro whether the epitope is delivered by pulsing from peptide, or expressed from within.
- CTL from a B7 donor displayed no reactivity to this epitope, although it had been immunodominant in another study Haas et al. [1996]

**HXB2 Location** Nef (128–137) **Author Location** Nef (128–137 SF2) **Epitope** TPGPGVRYPL **Immunogen** HIV-1 infection

Species (MHC) human (B7)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B7+ individuals that had a CTL response to this epitope broken down by group: 0/4 group 1, 0/3 group 2, and 1/1 group 3.

**HXB2 Location** Nef (128–137)

**Author Location** Nef (128–137)

**Epitope** TPGPGVRYPL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B7)

Keywords HIV exposed persistently seronegative

(HEPS), immunodominance

References Kaul et al. 2001a

- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused on different epitopes with HLA presenting molecules that have previously been associated with reduced risk of infection, and there was a shift in the response in the HEPS women upon late seroconversion to epitopes recognized by the HIV-1 infected women.
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B7 women, 4/5 HEPS and 5/6 HIV-1 infected women recognized this epitope.
- The dominant response to this HLA allele was to this epitope in 3 of the 4/5 HEPS cases and in 2 of the 5/6 HIV-1 infected women.
- Subject ML 1203 started with CTL responses to A\*6802 DTVLEDINL and to B7 FPVTPQVPLR prior to seroconversion, and upon seroconversion acquired additional responses to A\*6802 ETAYFILKL which became dominant, B7 TPGPG(V/I)RYPL, B7 IPRRIRQGL, and B7 SPRTLNAWV.

**HXB2 Location** Nef (128–137)

**Author Location** Nef (128–137)

Epitope TPGPGVRYPL

Immunogen HIV-1 infection

Species (MHC) human (B7)

References Appay et al. 2000

- Combined tetramer and intracellular cytokine staining was used to study the function of circulating CD8+ T cells specific for HIV and CMV.
- HIV-specific CD8+ T cells expressed lower levels of perforin than CMV-specific CD8+ T cells from the same donor, and this was associated with persistent CD27 expression on HIVspecific cells, suggesting impaired maturation.
- In most donors, between 50% and 95% of the activated virus-specific CD8+ T cells produced IFN- $\gamma$  and MIP-1 $\beta$  with a distinct subset that failed to produce TNF- $\alpha$

HXB2 Location Nef (128–137) Author Location Nef (128–137) Epitope TPGPGVPYPL

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Keywords** rate of progression, acute/early infection **References** Day *et al.* 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.
- Subjects with chronic HIV-1 infection recognized between 2-8 out of 11 B7-restricted epitopes.
- An acute seroconvertor homozygous for the B7 allele recognized five B7-restricted epitopes.
- The other acute seroconvertor failed to recognize any of the 11 B7-restricted epitopes tested.
- The B7-restricted CTL response was highly variable and there was no clearly dominant epitope.

**HXB2 Location** Nef (128–137)

Author Location Nef (128–137 BRU)

**Epitope** TPGPGVRYPL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B7)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- TPGPGVRYPL was recognized in 8/16 (50%) of individuals with HLA B7, and 1/9 (11%) of individuals with HLA B35. It was a high affinity HLA binder.

HXB2 Location Nef (128-137)

**Author Location** Nef

Epitope TPGPGVRYPL

Epitope name B7-TL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A3, B7, Cw7

**Keywords** dynamics, supervised treatment interruptions (STI), acute/early infection

References Yu et al. 2002a

- CTL responses in 18 acutely HIV-infected HLA-A3 (n=7) or -B7 (n=4) or both -A3 and B7 (n=7) positive individuals were studied.
- One individual, AC-06, was homozygous at all three class I alleles (A3, B7, Cw7), was treated during acute infection and had supervised treatment interruptions (STI). He had only two detectable CTL responses during acute infection, but after STI this broadened to 27 distinct epitopes including 15 restricted by HLA-A3, 11 by HLA-B7, and 1 by HLA-Cw7.
- 0/11 HLA-B7 individuals had detectable B7-restricted responses to this epitope during acute infection 10/15 of HLA-B7 epitopes tested were targeted by at least one person during acute infection. 2/4 individuals had detectable responses to this epitope after STI.

HXB2 Location Nef (128–137)

**Author Location** Nef

Epitope TPGPGVRYPL

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A2, A3, B7, Bw6

Keywords HAART, ART

References Appay et al. 2002

- Four HIV patients with prolonged clinically successful antiviral therapy but with ongoing evidence of replication and Nef mRNA transcription, showed specific T-cell responses by Elispot and Tetramer staining, maintained for 2-4 years after initiation of HAART.
- Nef epitope recognition was detected in all 4 subjects, gp120,
   Pol and Gag-specific in 1 or 2 subjects.
- The HIV-specific CD8+ T-cells had an intermediate maturation phenotype characterized by low levels of perforin and high levels of CD27 expression.

**HXB2 Location** Nef (128–137)

**Author Location Nef** 

Epitope TPGPGVRYPL

Subtype B, C

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade HIV component: p17 Gag, p24 Gag

Species (MHC) human, macaque (B7)

**Keywords** subtype comparisons, epitope processing, vaccine-specific epitope characteristics, im-

munodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the Aclade epidemic in Nairobi, Kenya. A DNA and MVA primeboost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

**HXB2 Location** Nef (128–137)

**Author Location** Nef

Epitope TPGPGVRYPL

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ

Keywords HIV exposed persistently seronegative

(HEPS)

References Koning et al. 2004

- A high-risk seronegative group of 29 homosexual men showed reduced cellular in vitro susceptibility for HIV infection and enhanced production of RANTES compared to 15 men who went on to seroconvert. Significantly higher frequencies of HLA A\*11, A\*31 and Cw\*15 were also found in the high risk seronegative men. Both groups of men had low frequencies of HIV-1 specific CD8+ T-cells, which may signify exposure more than protection from infection.
- No one, 0/9 HLA B7+ infection-resistant men, and 0/4 preseroconversion men who went on to become infected, reacted to this epitope.

**HXB2 Location** Nef (128–137)

**Author Location** Nef (126–135)

Epitope TPGPGVRYPL

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

 Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.  6/7 patients recognized this epitope, it was the most commonly recognized of 11 B\*07 epitopes.

**HXB2 Location** Nef (128–137)

Author Location (B consensus)

**Epitope** TPGPGVRYPL

Epitope name TL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Donor MHC A31, A68, B07, B70, Cw7, Cw1

Country United States.

Assay type Cytokine production, Intracellular cytokine

staining, Chromium-release assay, Flow cyto-

metric T-cell cytokine assay

Keywords assay standardization/improvement, memory

cells, characterizing CD8+ T cells

References Lichterfeld et al. 2004c

- Using a flow-cytometric cytotoxicity assay based on caspase-3
  activation in dying target cells, it was shown that the subset of
  HIV-1-specific CD8+ T cells secreting both IFN-gamma and
  TNF-alpha exhibit stronger cytotoxic activity than those secreting only IFN-gamma. These cells also exhibited stronger intracellular perforin expression. No association between HIV-1-specific CD8+ T-cell maturation phenotypes and intracellular
  perforin expression was found.
- 1/9 individuals recognized this epitope.

**HXB2 Location** Nef (128–137)

Author Location Nef

Epitope TPGPGVRYPL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Country United Kingdom.

Assay type Tetramer binding, T-cell Elispot, Intracellular

cytokine staining

Keywords rate of progression, acute/early infection, char-

acterizing CD8+ T cells, immune dysfunction

References Papagno et al. 2004

Acute HIV-1 infection induces massive activation of HIV-specific and non-HIV-specific CD8+ T-cells resulting in differentiation of these cells. High differentiation of CD8+ T-cells is correlated with disease progression. Differentiation is a natural process but it can be driven by elevated immune activation, such as in HIV infection.

HXB2 Location Nef (128–137)

**Author Location** Nef (128–137)

Epitope TPGPGVRYPL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

**Species (MHC)** human (B7 supertype)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization

## References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

**HXB2 Location** Nef (128–137)

**Author Location** Nef (128–137 subtype B)

Epitope TPGPGVRYPL

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B7, B\*8101)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix systemic CD8+ T cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

**HXB2 Location** Nef (128–137)

**Author Location** Nef (subtype B)

Epitope TPGPGVRYPL

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B7, B\*8101)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- Clade A version of the epitope: TPGPGIRYPL, clade D version: TPGPGIRYPL.

HXB2 Location Nef (128–137)

Author Location Nef

Epitope TPGPGIRYPL

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted – 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.

- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was recognized by 1/22 HEPS control sex workers, ML851.

**HXB2 Location** Nef (130–139)

Author Location Nef (130–139 BRU)

Epitope GPGVRYPLTF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- GPGVRYPLTF was recognized in 0/10 (0%) of individuals with HLA B7, and 1/11 (9%) of individuals with HLA B35, although it was a high affinity HLA binder.

**HXB2 Location** Nef (130–143)

**Author Location** Nef (130–143 LAI)

Epitope GPGVRYPLTFGWCY

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B\*57)

References Goulder et al. 1996b

- CTL response to this epitope observed in 4 long-term survivors.
- Peptide defined on the basis of B\*5801 binding motif, yet not cross-restricted except at high concentrations.

**HXB2 Location** Nef (130–143)

**Author Location** Nef (121–141)

Epitope GPGVRYPLTFGWCY

Immunogen HIV-1 infection

Species (MHC) human (B57)

References Ferrari et al. 2000

• One of the 51 HIV-1 epitopes selected by Ferrari *et al.* as good candidate CTL epitopes for vaccines by virtue of being conserved and presented by common HLA alleles.

**HXB2 Location** Nef (130–143)

Author Location Nef (128-141)

Epitope GPGVRYPLTFGWCY

Immunogen HIV-1 infection

Species (MHC) human (B57)

Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 7/7 patients recognized this epitope.

**HXB2 Location** Nef (130–144)

Author Location Nef (130–144 HXB2)

Epitope GPGVRYPLTFGWCYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type T-cell Elispot

Keywords supervised treatment interruptions (STI), im-

munodominance, early treatment

References Addo et al. 2003

- Comprehensive Elispot screening of 504 overlapping peptides that spanned all HIV-1 proteins in 57 HIV-1 infected individuals. Four groups of patients were compared: 23 were untreated; 12 patients were chronically infected and treated; 22 started treatment during acute infection, 11 continuously treated and 11 with STI.
- 63% of the peptides were recognized the most frequent responses were directed against Nef (95%) and p24-Gag (88%). p17 was the most frequently recognized protein after correction for protein length. A median of 18 peptides (range 2-42) were recognized per person. The most frequently (>20%) recognized peptides were located in conserved regions within clade B sequences. Vpu was rarely recognized.
- A decrease in the total magnitude and in the breadth of CTL responses was detected in patients with treated acute infection in comparison to chronically infected treated or untreated individuals. No correlation between plasma viral load and HIV-1 specific T-cell responses was observed.
- The authors did not note the reference strain, but based on the peptide sequences provided it appears to be HXB2.
- Responses to this peptide were detected in 24% of the study subjects, and it was one of the 25 most frequently recognized peptides.

**HXB2 Location** Nef (132–144)

**Author Location** Nef

Epitope GIRYPLTFGWCFK

Immunogen

Species (MHC) human

Keywords subtype comparisons

References Jubier-Maurin et al. 1999

- 41 new HIV-1 strains describing envelope subtypes of HIV-1
   A-H were genetically characterized in the nef region 34 subtypes were classified in the same subtype in nef and env and 7 of the 41 strains were recombinants.
- This region was defined as a CTL epitope region that is conserved among HIV-1 M group subtypes.

HXB2 Location Nef (132–147)

Author Location Nef (132–147 BRU)

**Epitope** GVRYPLTFGWCYKLVP **Immunogen** HIV-1 infection

Species (MHC) human (A1, B8)

References Hadida et al. 1992

• HIV-1 specific CTLs detected in lymphoid organs.

**HXB2 Location** Nef (132–147)

**Author Location** Nef (132–147 BRU)

Epitope GVRYPLTFGWCYKLVP

Immunogen HIV-1 infection

Species (MHC) human (B18)

References Culmann et al. 1991

• Nef CTL clones from HIV+ donors.

**HXB2 Location** Nef (132–147)

Author Location Nef (132-147)

Epitope GVRYPLTFGWCYKLVP

Immunogen vaccine

*Vector/Type:* DNA, DNA with protein boost *Strain:* B clade LAI *HIV component:* Gag,

Nef. Tat Adjuvant: IL-18

Species (MHC) mouse (H-2<sup>d</sup>)

Keywords Th1

References Billaut-Mulot et al. 2001

- DNA vaccinated BALB/c mice primed and boosted with the multiepitopic vaccine with IL18 showed lymphoproliferative responses 7 weeks post immunization.
- Strong but non-lasting HIV-specific CTL responses were detected by a Cr-release assay and DNA prime/DNA boost was more effective than DNA prime protein boost.
- Immunization with either the multiepitopic DNA or with the mixed DNA vaccine induced HIV-1 specific Th1 cytokines (IL-2 and IFN-gamma)
- Co-administration of IL18 increased T-cell responses but decreased anti-HIV antibody levels.

**HXB2 Location** Nef (133–141)

**Author Location** Nef (133–141)

Epitope TRYPLTFGW

Immunogen HIV-1 infection

Species (MHC) human (A33)

Keywords optimal epitope

**References** Frahm *et al.* 2007

**HXB2 Location** Nef (133–148)

**Author Location** Nef (133–148 LAI)

Epitope VRYPLTFGWCYKLVPV

Subtype B

**Immunogen** 

Species (MHC) human (B57)

**References** Brander & Walker 1996

• P. Goulder, pers. comm.

**HXB2 Location** Nef (134–141)

Author Location (C consensus)

**Epitope RYPLTFGW** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** characterizing CD8+ T cells

## References Kiepiela et al. 2004

- HLA class I restricted CD8+ T-cell responses against HIV-1
  were analyzed in African patients. Significantly more responses
  were shown to be HLA-B restricted. Viral load, CD4 count,
  and thus rate of disease progression were also associated with
  HLA-B alleles. In addition, the selection pressure imposed on
  HIV-1 by HLA-B alleles was shown to be substantially greater
  than by other alleles.
- This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** Nef (134–141)

Author Location (C consensus)

**Epitope** RYPLTFGW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords epitope processing, rate of progression, opti-

mal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed.
   Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- Mutational patterns in a residue outside of the optimized epitope of RYPLTFGW are associated with the presence of the HLA presenting molecule in the host.

**HXB2 Location** Nef (134–141)

Author Location Nef (138-147 LAI)

Epitope RYPLTFGW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*2402 epitope.

**HXB2 Location** Nef (134–141)

**Author Location** Nef (169–176)

**Epitope** RYPLTFGW

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

Assay type Other

Keywords HLA associated polymorphism

References Boutwell & Essex 2007

 All HIV-1 subtype C protein sequences were analysed, and 94 HLA-associated amino acid polymorphisms were found that were well distributed throughout the proteome. These sequences were compiled from publicly available databases and originated in South Africa and Botswana. Many polymorphisms were associated with multiple HLA allele classes. 12% of negative associations were found as well, where there was an association of virus expression of the consensus amino acid with a specific HLA allele. HLA-B alleles were involved in most of the associations. 19 HLA-associated polymorphisms were embedded in previously defined epitopes presented by an appropriate HLA restricting molecule.

 RYPLTFGW was a previously defined A\*2402 presented epitope that encompassed an A\*24 associated polymorphism, Ry-PLTFGW.in the second position.

**HXB2 Location** Nef (134–141)

**Author Location** Nef (138–147 SF2)

Epitope RYPLTFGW

Immunogen HIV-1 infection

Species (MHC) human (A24)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A24+ individuals that had a CTL response to this epitope broken down by group: 0/0 group 1, 2/3 group 2, and 0/0 group 3.

**HXB2 Location** Nef (134–141)

**Author Location** Nef

Epitope RYPLTFGW

**Epitope name** A24-RW8(Nef)

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A24)

**Donor MHC** A24, B7, B27

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.

- Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals. Patient C displayed the greatest response to B27-KK10(p24), and in decreasing order also responded to A24-RW8(Nef), B7-IL9(gp41), A24-RL9(gp41), A24-YL8(gp41), and B7-TM9(Nef).

**HXB2 Location** Nef (134–141)

Author Location Nef (134-141)

Epitope RYPLTFGW

Subtype B

**Immunogen** HIV-1 infection **Species** (MHC) human (A33)

Donor MHC A3, A33, B14, B35, Cw\*0401, Cw\*0802

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** binding affinity, acute/early infection, earlyexpressed proteins

References Cao et al. 2003

- All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.
- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

**HXB2 Location** Nef (134–141)

Author Location Nef (134-141 LAI)

 ${\bf Epitope} \ \ {\sf RYPLTFGW}$ 

Subtype B

Immunogen

Species (MHC) human (B27)

References Culmann 1998

· Optimal peptide defined by titration.

**HXB2 Location** Nef (134–143)

Author Location Nef (134–143)

Epitope RYPLTFGWCF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*24)

**Donor MHC** A\*03, A\*24, B\*35, B\*40

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** acute/early infection, variant crossrecognition or cross-neutralization, superinfection, characterizing CD8+ T cells

References Yang et al. 2005b

- An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- The response to the peptide that carried this epitope, RYPLTFG-WCF, was present before superinfection but waned afterward.
   The epitope from the second strain had a mutation, ryplCfgwcf.
   A second overlapping epitope in the reactive peptide might be involved, the B\*35 epitope YPLTFGWCF.

**HXB2 Location** Nef (134–143)

Author Location Nef (138–147 SF2)

Epitope RYPLTFGWCF

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

References Ikeda-Moore et al. 1997

- Defined using reverse immunogenetics 59 HLA-A\*2402 binding peptides were predicted by searching for A\*2402 anchors in HIV proteins (Tyr at 2, and Phe, Leu or Ile at the C term) 53 of the 59 peptides bound A\*2402.
- This peptide induced CTL in 3/4 HIV-1 + people tested.
- RYPLTFGWCF bound to A\*2402 strongly, the epitope can be processed in a vaccinia construct and presented – two specific CTL clones were obtained.

**HXB2 Location** Nef (134–143)

**Author Location** Nef (138–147)

Epitope RYPLTFGWCF

Epitope name Nef138-10

Subtype B

Immunogen vaccine

Vector/Type: Sendai virus vector system (SeV)

Species (MHC) human (A\*2402)

References Kawana-Tachikawa et al. 2002

- A Sendai virus vector system (SeV) was developed that expressed HLA-A\*2402-restricted class I/peptide complexes; this system could be used to detect responses and has the potential to elicit immune responses.
- MHC class I/peptide tetramers could be made using this system that bound to epitope-specific CTLs in PBMCs.
- Cells transfection with SeV modified to express A\*2402-HIV epitope complexes induced CTL mediated specific cell lysis.

**HXB2 Location** Nef (134–143)

**Author Location** Nef

Epitope RYPLTFGWCF

**Epitope name** Nef138-10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*2402)

Donor MHC A\*2402

Country Japan.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay

Keywords binding affinity, epitope processing, immun-

odominance, escape

References Furutsuki et al. 2004

• 70% of Japanese people carry HLA A\*2402, and the rFpltfgwcf (2F) escape variant of this A\*2402 epitope was found to be positively selected in Japan; reversion to wild-type in HLA-A24 negative individuals occurred very slowly over years. The 2F escape variant appears to be common in Japan due to escape and then transmission of this form in the population. The mechanism of escape appeared to be in processing of Nef and antigen presentation rather than HLA binding since both wild-type and 2F variant bound to HLA-A\*2402 with almost same efficiency; the authors suggest the epitope may be cleaved at position 5 with a higher frequency when the 2F mutation is present.

**HXB2 Location** Nef (134–143)

**Author Location** Nef (134–143 BRU)

Epitope RYPLTFGWCY

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (A24)

**Keywords** binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- RYPLTFGWCY was recognized in 5/12 (42%) of individuals with HLA A24. It was a moderate affinity HLA-A24 binder.

**HXB2 Location** Nef (134–143)

**Author Location** Nef (134–143)

Epitope RYPLTFGWCY

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (A24)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

Keywords vaccine-induced epitopes, variant cross-

recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

**HXB2 Location** Nef (134–143)

**Author Location** Nef (134–143 HXB2)

Epitope RYPLTFGWCY
Subtype B, CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A24)
Country Viet Nam.

Assay type HLA binding

**Keywords** subtype comparisons, computational epitope prediction, variant cross-recognition or cross-

neutralization, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-A11, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- CRF01\_AE variant ryplCfgwcy had same HLA-binding score as the HXB2 epitope.

**HXB2 Location** Nef (134–143)

**Author Location** Nef

Epitope RYPLTFGWCF Immunogen HIV-1 infection Species (MHC) human (A24)

**Donor MHC** A\*24, A\*32, B\*07, B\*18, Cw\*07

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, characterizing CD8+ T cells,

reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs.
   Escape variants were commonly detected in maternal plasma.
   Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- Variant rFpltfgwcf was detected in increasing frequencies in clones from an A24+ infant, but was absent in all sequences from the A24- mother at delivery, revealing selective pressure as early as 3 months of age.

**HXB2 Location** Nef (134–143)

**Author Location** Nef (134–143 BRU)

Epitope RYPLTFGWCY Subtype B, CRF02\_AG Immunogen HIV-1 infection Species (MHC) human (A24, B35)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFNγ **Keywords** subtype comparisons References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 3/9 CRF02\_AG-infected Ivorians, and 1/9 B-infected French subjects.
- A C-term F was most common in both the CRF02 and B clade infected subjects, and subjects that carried the F, RYPLTFG-WCf, reacted with the peptide. One Ivorian that recognized the peptide carried the form RfPLTFGWCf.

**HXB2 Location** Nef (134–144)

Author Location Nef (134-144 LAI)

Epitope RYPLTFGWCYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B18)

Keywords review, escape

References Couillin et al. 1994; Goulder et al. 1997a

- · Mutational variation in HIV epitopes in individuals with appropriate HLA types can result in evasion of CTL response.
- Goulder et al. [1997a] is a review of immune escape that summarizes this study.

**HXB2 Location** Nef (134–144)

**Author Location** Nef (134–144)

Epitope RYPLTFGWCYK

**Epitope name** RYP

Immunogen HIV-1 infection Species (MHC) human (B18)

Keywords HAART, ART, acute/early infection

References Oxenius et al. 2000

- Patients who started therapy at acute HIV-1 infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load - three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- None of the 8 study subjects recognized this epitope but none were HLA B18+

**HXB2 Location** Nef (135–143)

Author Location p17

Epitope YPLTFGWCF Immunogen HIV-1 infection Species (MHC) human (A2)

Assay type Intracellular cytokine staining

Keywords immunodominance, genital and mucosal immunity

References Kaul et al. 2003

- Predefined immunodominant peptide responses were used to compare CD8+ T cells responses in the blood and cervix of 16 HIV+ Kenyan sex workers. Cervical responses were detected in 8/10 women from whom adequate samples could be obtained. The frequency of the CD8+ T cell response in the genital tract was comparable to the blood, with a trend toward being slightly higher.
- The immunodominant response was to this epitope in the PBMC of 1/16 patients (Kaul et al. 2001, AIDS, 107:1303).

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143 LAI)

**Epitope** YPLTFGWCY

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (B\*0702)

Keywords epitope processing

References Lucchiari-Hartz et al. 2000

- Five naturally processed MHC class I ligands were identified in Nef in the conserved immunogenic region Nef between
- All five could be transported by TAP, and 4/5 had N-termini that were major cleavage points for the proteasome, only one had extended precursor fragments.
- YPLTFGWCY is the naturally processed ligand for B7, and this epitope is the only one of the five that may require trimming at the N-termini.
- YPLTFGWCY is present in low copy number in the cell, possibly due to a predominant proteasomal cleavage site between Y and P.

**HXB2 Location** Nef (135–143)

**Author Location** (C consensus)

**Epitope** YPLTFGWCF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*18)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- YPLTFGWCF is an optimal epitope for B\*5301, B\*18, and

**HXB2 Location** Nef (135–143)

Author Location Nef (135–143 LAI)

**Epitope** YPLTFGWCY

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B\*1801)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is a B\*1801 epitope.

HXB2 Location Nef (135-143)

**Author Location** Nef (135–143 HXB2)

**Epitope** YPLTFGWCF

Epitope name YF9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*1801)

**Donor MHC** A\*0201, A\*2501, B\*1801, B\*5101, • YPLTFGWCF is an optimal epitope for B\*5301, B\*18, and

Cw\*0102, Cw\*1203

Assay type CD8 T-cell Elispot - IFNγ

Keywords escape, immune evasion, optimal epitope, HIV-1

References Liu et al. 2006

• T-cell immunity and the impact of selection were longitudinally studied in a patient from the time of transmission. New epitopes were detected at sites that were shown to be under positive selection. Env was found to be the major target of CTLs and most of the new epitopes were found here. One third of the sites examined were associated with epitope escape mutations from CTL. Acquisition of mutations found at high frequency in the database was suggested to be a result of reversion of CTL epitopes in the absence of immune targeting.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143)

**Epitope** YPLTFGWCF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*35)

**Donor MHC** A\*03, A\*24, B\*35, B\*40

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** acute/early infection, variant cross-

recognition or cross-neutralization, superin-

fection

References Yang et al. 2005b

- An individual infected with a drug-resistant strain of HIV-1 subtype B with controlled viremia became superinfected with another subtype B strain. The second strain outgrew the first despite lower replication capacity and the same viral phenotype. The strains showed differences in their epitope sequences. The CTL responses to the first strain declined after superinfection, followed by some adaptation of targeting to the new epitopes of the second strain. Differences in the recognized epitopes were suggested to have contributed to the poor immune containment of the second strain.
- The response to the peptide that carried this epitope, YPLTFG-WCF, was present before superinfection but waned afterward. The epitope from the second strain had a mutation, yplCfgwcf. A second overlapping epitope in the reactive peptide might be involved, the A\*24 epitope RYPLTFGWCF.

**HXB2 Location** Nef (135–143)

**Author Location** (C consensus)

**Epitope** YPLTFGWCF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*35)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- B\*35.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143)

**Epitope** YPLTFGWCY

Immunogen

Species (MHC) human (B\*5301)

Keywords optimal epitope

References Frahm et al. 2007

**HXB2 Location** Nef (135–143)

**Author Location** (C consensus)

**Epitope** YPLTFGWCF

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (B\*5301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords rate of progression, optimal epitope

References Kiepiela et al. 2007

- A comprehensive analysis of 160 class I T cell responses in 578 individuals from KwaZulu-Natal, South Africa was performed. Gag-specific responses were associated with lowering viremia, while Env, accessory and regulatory protein-specific responses were associated with higher viremia.
- YPLTFGWCF is an optimal epitope for B\*5301, B\*18, and B\*35.

**HXB2 Location** Nef (135–143)

**Author Location** 

Epitope YPLTFGWCY

Epitope name Nef-YY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B\*5301, B35)

Donor MHC A\*3002 A\*3201 B\*4501 B\*5301 Cw\*0401 Cw\*1202

Keywords HAART, ART

References Sabbaj et al. 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.

- Subject 00RCH33 was on HAART had a viral load of 2900 and CD4 count of 727 and also recognized the epitopes HIGPGRAFY, gp160(310-318), HLA A\*3002; AETFYVDGA, RT(437-445), HLA B\*4501; and RSLYNTVATLY, p17(76-86), HLA A\*3002.
- Among HIV+ individuals who carried HLA B53, 8/15 (53%) recognized this epitope – one subject also carried B7, previously shown to restrict this epitope.
- Among HIV+ individuals who carried HLA B35, 13/19 (68%) recognized this epitope.

HXB2 Location Nef (135–143) Author Location Nef (subtype D) Epitope YPLTFGWCF

Subtype D

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B18)

References Kaul et al. 2000

- 11/16 heavily HIV exposed but persistently seronegative sexworkers in Nairobi had HIV-specific CD8 gamma-IFN responses in the cervix systemic CD8+ T cell responses tended to be to the same epitopes but at generally lower levels than cervical CD8+ T cell responses.
- Low risk individuals did not have such CD8+ cells.
- CD8+ T cell epitopes: DTVLEDINL (3 individuals), SLYNT-VATL (4 individuals), LSPRTLNAW (3 individuals) and YPLT-FGWCF (4 individuals) were most commonly recognized by the HIV-resistant women.

HXB2 Location Nef (135–143) Author Location Nef (135–143 LAI)

**Epitope** YPLTFGWCY

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B18)

**References** Culmann *et al.* 1991; Culmann-Penciolelli *et al.* 1994

• Nef CTL clones from HIV+ donors.

**HXB2 Location** Nef (135–143)

Author Location Nef (135–143 SF2)

Epitope YPLTFGWCY Immunogen HIV-1 infection Species (MHC) human (B18)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.

• Number of HLA-B18+ individuals that had a CTL response to this epitope broken down by group: 0/3 group 1, 1/2 group 2, and 0/0 group 3.

**HXB2 Location** Nef (135–143)

**Author Location Nef** 

Epitope YPLTFGWCF

Immunogen HIV-1 infection

Species (MHC) human (B18)

**Keywords** HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2002

- Neisseria gonorrhea cervititis in 9 HIV+ Kenyan sex workers caused a functional deficiency in IFN-gamma production in HIV-1 epitope-specific CD8+ T-cells, detected by intracellular cytokine production and tetramer assays, while not affecting the total number of epitope-specific CTLs.
- Ghonorrhea caused the weaker HIV-1 specific CTL responses in 4 HIV-1 exposed persistently seronegative (HEPS) women to become undetectable by Elispot and tetramer assays, and CMV-specific CTL in 2 HEPS subjects were shown to have impaired function with regard to IFN-gamma production.

**HXB2 Location** Nef (135–143)

**Author Location Nef** 

Epitope YPLTFGWCY

Epitope name B18-YY9(Nef)

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B18)

**Donor MHC** A30, A32, B18, B27

**Keywords** HAART, ART, supervised treatment interruptions (STI)

References Altfeld et al. 2002b

- Peripheral blood (PB) and lymph node (LN) CD8+ T-cell responses were compared in 15 asymptomatic HIV-1 infected patients using all known optimal CTL epitopes (http://hiv-web.lanl.gov/content/hiv-db/REVIEWS/brander2001.html) for each person's class I HLA alleles.
- 60 epitope responses were detected in both PB and LN samples
  of the 15 patients, and an additional 8 responses were detected
  only in LN. The total magnitude of the response was similar
  in LN and PB, but the percentage of CD8+ T cells in the LN
  is lower so the number of HIV-specific cells per million CD8+
  T-cells is higher in the LN.
- 1 year post-HAART treatment in five patients studied, the magnitude of the CD8 T-cell response was decreased in both LN and PB, but more dramatically in PB, and 13/25 epitope responses in the PB became undetectable, in contrast to 5/26 in the LN.
- Treatment interruption following HAART induced resulted in increased viremia accompanied by the restoration of the detection of 13 epitopes that had become undetectable in the PB, and the addition of 9 novel epitope responses.
- Breakdowns of epitope responses were shown for 4 individuals.
   Patient D displayed the greatest response to B27-KK10 (p24), and also responded to A30-RY11(p17), A32-PW10(RT), A30-KY11(RT), A32-RW10(gp120), and B18-YY9(Nef).

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143)

Epitope YPLTFGWCY

Epitope name YY9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B18)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Flow cytometric T-cell cytokine

assay

**Keywords** optimal epitope **References** Allen *et al.* 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- This epitope did not vary.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143)

**Epitope** YPLTFGWCY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B18)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** Nef (135–143)

**Author Location** (C consensus)

**Epitope** YPLTFGWCF

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** human (B18, B\*5301)

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

Keywords cross-presentation by different HLA, charac-

terizing CD8+ T cells

References Kiepiela et al. 2004

HLA class I restricted CD8+ T-cell responses against HIV-1
were analyzed in African patients. Significantly more responses
were shown to be HLA-B restricted. Viral load, CD4 count,
and thus rate of disease progression were also associated with
HLA-B alleles. In addition, the selection pressure imposed on
HIV-1 by HLA-B alleles was shown to be substantially greater
than by other alleles.

 This epitope was suggested to be the epitope within a longer reactive peptide based on correspondence with a known epitope in the HIV database. Also, a significantly higher frequency of people in the Durban cohort who reacted with the peptide had this HLA type.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143)

**Epitope** YPLTFGWCY

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (B18, B49)

Keywords HIV exposed persistently seronegative

(HEPS), immunodominance

References Kaul et al. 2001a

- Variants YPLTFGWC(Y/F) are specific for the B/D clades.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.
- Responses in HEPS women tended to be lower, and focused
  on different epitopes with HLA presenting molecules that have
  previously been associated with reduced risk of infection, and
  there was a shift in the response in the HEPS women upon late
  seroconversion to epitopes recognized by the HIV-1 infected
  women
- 43/91 HEPS women had CD8+ responses and detection of HIV-1-specific CTL in HEPS women increased with the duration of viral exposure.
- Among HLA-B18 women, 1/4 HEPS and 8/9 HIV-1 infected women recognized this epitope, likelihood ratio 5.3, p value 0.04, and HEPS women tended to respond to FRDYV-DRF(Y/F)K, while infected women tended to respond to YPLT-FGWC(Y/F)
- The dominant response to this HLA allele was to this epitope for the one reactive HEPS case and in all 8/9 HIV-1 infected women.
- Differences in epitope specificity were only seen for responses restricted by class I HLA alleles A2, A24, A\*6802, B14, and B18, previously shown to be associated with resistance to HIV-1 in this cohort.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (139–147 SF2)

Epitope YPLTFGWCF

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Shiga et al. 1996

• Binds HLA-B\*3501.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143 BRU)

Epitope YPLTFGWCY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- YPLTFGWCY was recognized in 2/13 (15%) of individuals with HLA B7, and 11/14 (79%) of individuals with HLA B35, and it was a moderate affinity HLA binder.

**HXB2 Location** Nef (135–143)

**Author Location Nef** 

**Epitope** YPLTFGWCY

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35)

**Donor MHC** A3, A11, B35, B51

References Sabbaj et al. 2002

Keywords mother-to-infant transmission

- IFNgamma T-cell responses in breast milk of 5 HIV-1 infected women from the US and 6 from Zambia were tested with using Elispot. 11/11 women responded to Gag, 8/11 responded to Pol, 7/11 women to Nef, and 2/5 women to Env peptide pools. These responses were shown to be primarily due to CD8+ T-cells in one woman, and another woman had cytolytic responses measured by Cr-release.
- T-cells in breast milk from a volunteer who was HLA A3, A11, B35, B51 induced IFNgamma after stimulation with a peptide that carries known B35 epitope YPLTFGWCY.
- The frequencies of responses in the two compartments differed, and 2/4 women that responded to epitopes in Nef 101-205 and Pol 601-710 showed responses in breast milk but no detectable responses in peripheral blood cells.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143)

**Epitope** YPLTFGWCY

Epitope name YY9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B35)

Donor MHC A\*0201, A\*0301, B\*3501, B\*51, Cw\*04,

Cw\*06

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Chromium-release assay

Keywords escape, acute/early infection

References Bansal et al. 2005

• Patients with acute or early infection were shown to preferentially target variable peptides with higher entropy while those with chronic infection showed responses towards more conserved peptides with lower entropy. In longitudinally followed subjects, responses to variable proteins declined over time while responses to conserved proteins increased. The decline is suggested to be due to CTL escape.

over time.

**HXB2 Location** Nef (135–143)

**Author Location Nef** 

Epitope YPLTFGWCY

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B49)

Keywords subtype comparisons, HIV exposed persistently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- · A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades - such crossreactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A subtype consensus is identical to the B clade epitope.
- The D subtype consensus is ypltfgwcF.

**HXB2 Location** Nef (135–143)

Author Location Nef (subtype B)

**Epitope** YLPTFGWCY

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (B49)

**Keywords** subtype comparisons

References Rowland-Jones et al. 1998b

- · HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi – these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found - B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- This epitope is conserved among A and B clade viruses.
- The Clade D version of the epitope, YPLTFGWCF, was preferentially recognized by CTL.

**HXB2 Location** Nef (135–143)

**Author Location** 

Epitope YPLTFGWCY

Immunogen HIV-1 infection Species (MHC) human (B49)

Keywords HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001c

- · This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted - 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope, YPLTFGWC(Y/F), was recognized in 1/22 HEPS sex worker controls (ML1668)

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143)

**Epitope** YPLTFGWCF

Immunogen

Species (MHC) human (B53)

Keywords optimal epitope

**References** Frahm *et al.* 2007

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143 BRU)

**Epitope** YPLTFGWCY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- YPLTFGWCY was recognized in 2/13 (15%) of individuals with HLA B7, and 11/14 (79%) of individuals with HLA B35, and it was a moderate affinity HLA binder.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143)

**Epitope** YPLTFGWCY

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (B7 supertype)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

Keywords vaccine-induced epitopes, variant cross-

recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

• After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the most frequently recognized of the 31 epitopes that the vaccinated volunteers responded to.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143)

**Epitope** YPLTFGWCY

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (B7, B18, B35, B49, B53)

**Donor MHC** A3, A31, B18, B39; A1, A3, B8, B35

Country United States.

Assay type Intracellular cytokine staining, Flow cytomet-

ric T-cell cytokine assay

Keywords HAART, ART, escape, variant cross-

recognition or cross-neutralization

References Casazza et al. 2005

• Evidence of continued viral evolution during HAART therapy despite low viral load was found in 1/5 patients studied.

• A predominant sequence of ypltfgwcF was found in 2 patients (in 11/11 and 15/15 sequences) that cross-recognized the peptide used for screening, YPLTFGWCY. In patient B, CD8 T-cell response of 0.95% was found for the dominant variant, while the response for the screening epitope ypltfgwcy was 0.58%. In patient F, the frequency of response did not differ significantly between the 2 variants. Assays for both patients were done immediately prior to the initiation of therapy.

**HXB2 Location** Nef (135–143)

**Author Location** Nef (135–143 BRU)

Epitope YPLTFGWCY

Subtype B, CRF02\_AG

Immunogen HIV-1 infection

Species (MHC) human (B7, B18, B53)

Country Cote D'Ivoire.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 3/9 CRF02\_AG-infected Ivorians, and 0/9 B-infected French subjects.
- The three Ivorians that recognized this peptide carried three different variants of the epitope: identical to LAI: YPLTFG-WCY, YPLTFGWCf, and fPLTFGWCf. 6/8 Ivorians carried a variant, 5/5 B clade infections were not identical.

**HXB2 Location** Nef (136–144)

Author Location Nef (136–144 BRU)

Epitope PLTFGWCYK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

**Keywords** binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- PLTFGWCYK was recognized in 3/12 (25%) of individuals with HLA A3. It was a low affinity HLA-A3 binder.

**HXB2 Location** Nef (136–145)

**Author Location** Nef (136–145)

Epitope PLTFGWCYKL

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Keywords binding affinity, dendritic cells, Th1

References Wilson et al. 1999b

- Dendritic cells are the most potent for priming T cell responses
   DCs can stimulate autologous CTL responses from T cells cultured from HIV negative donors.
- Th1-biasing cytokines IL-12 or IFN alpha enhance CTL responses *in vitro* whether the epitope is delivered by pulsing from peptide, or expressed from within.
- B7 and A2 Nef epitopes were studied and the relative binding affinity of A2 epitopes for A2 was: PLTFGWCYKL greater than VLEWRFDSRL which was much greater than AFHH-VAREL.
- Noted in Brander et al., 1999 this database, to be A\*0201.

**HXB2 Location** Nef (136–145)

**Author Location** Nef (136–145 LAI)

Epitope PLTFGWCYKL

Subtype B

Immunogen

Species (MHC) human (A\*0201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0201 epitope.

**HXB2 Location** Nef (136–145)

Author Location Nef (136–145 LAI)

Epitope PLTFGWCYKL

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Keywords epitope processing

References Lucchiari-Hartz et al. 2000

- Five naturally processed MHC class I ligands were identified in Nef in the conserved immunogenic region Nef between 123-152.
- All five could be transported by TAP, and 4/5 had N-termini that were major cleavage points for the proteasome, only one had extended precursor fragments.
- The CTL that recognized PLTFGWCYKL also recognized PLTFGWCYKLV, and both forms of the epitope are naturally processed and both seem to be the direct product of a proteasomal digest, although in low copy number.

**HXB2 Location** Nef (136–145)

**Author Location** Nef (136–145)

Epitope PLTFGWCYKL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot •

granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- Three of seven patients responded to this peptide with GzB producing cells and three of the patients responded with IFNgamma producing cells. Only one patient had both a GzB and IFN-gamma response.

**HXB2 Location** Nef (136–145)

**Author Location Nef** 

Epitope PLTFGWCFKL

Epitope name P10L Immunogen vaccine

Vector/Type: measles virus (MV) Strain: multiple epitope immunogen HIV compo-

*nent:* gp140, gp140∆V3

Species (MHC) transgenic mouse (A\*0201)

Assay type Chromium-release assay, Flow cytometric T-

cell cytokine assay

**Keywords** memory cells, vaccine antigen design, antibody generation, characterizing CD8+ T cells

References Lorin et al. 2005

A recombinant measles MVSchw virus expressing an HIV-1derived polyepitope effectively primed HLA-A\*0201-restricted
CTL responses against multiple conserved HIV-1 epitopes in
HLA-A\*0201 transgenic mice. Also, a recombinant MVSchw
virus expressing gp140 with deleted V1, V2, and V3 loops
successfully induces neutralizing antibodies against HIV-1. A
live attenuated measles vaccine could provide a safe and efficient pediatric vaccination vector for simultaneous vaccination
against HIV and measles.

**HXB2 Location** Nef (136–145)

Author Location Nef (136–145)

Epitope PLTFGWCFKL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords subtype comparisons

References Durali et al. 1998

- Cross-clade CTL response was studied by determining the CTL activity in seven patients from Bangui, (6 A subtype, and 1 AG recombinant infections) and one A subtype infection from a person living in France originally from Togo, to different antigens expressed in vaccinia.
- Pol reactivity: 8/8 had CTL to A subtype, and 7/8 to B subtype, and HIV-2 Pol was not tested.
- Gag reactivity: 7/8 reacted with A or B subtype gag, 3/8 with HIV-2 Gag.
- Nef reactivity: 7/8 reacted with A subtype, and 5/8 with B subtype, none with HIV-2 Nef.
- Env reactivity: 3/8 reacted with A subtype, 1/8 with B subtype, none with HIV-2 Env.
- Patient B18 had the greatest breadth and diversity of response, and recognized Gag SLYNTVATL and Nef PLTFGWCFKL.

HXB2 Location Nef (136–145) Author Location Nef (157–166) Epitope PLTFGWCFKL Immunogen vaccine

Vector/Type: DNA prime with vaccinia boost

Species (MHC) human (A2)

References Woodberry et al. 1999

- A polyepitope vaccine was generated in a vaccinia construct that contiguously encoded seven epitopes, all presented by HLA A-2.
- HHD mice have a transgene of HLA A2 linked to the transmembrane and cytotoxic domains of H-2D<sup>d</sup> – this transgene is the only MHC molecule expressed in the mice.
- CTL responses to Gag (77-85) SLYNTVATL, Pol (476-484) ILKEPVHGV, gp120 (120-128) KLTPLCVTL, and Nef (190-198) AFHHVAREL were observed in HIV polytope HHD-vaccinated mice, and these responses were enhanced with vaccinia boost.
- No CTL immune responses were generated against HLA A2restricted HIV epitopes Nef 157-166 (PLTFGWCYKL), Pol 346-354 (VIYQYMDDL), and Nef 180-189 (VLEWRFDSRL)
- Sixteen HLA A2+ patients were tested for their ability to make CTL responses by peptide restimulation in culture with the epitopes selected for inclusion in the polytope – one individual recognized all seven of these epitopes; 7 patients had CTL cultures able to recognize at least one of the epitopes, and 6 of those 7 recognized more than one epitope, but they were not able to test all peptides for all patients; many patients only had three peptides tested.
- PLTFGWCFKL was recognized by 1 of the HLA-A2 patients.

**HXB2 Location** Nef (136–145)

**Author Location** Nef (135–144 93TH253 subtype CRF01)

Epitope PLTFGWCYKL Subtype CRF01\_AE Immunogen HIV-1 infection Species (MHC) human (A2)

**Keywords** subtype comparisons **References** Bond *et al.* 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- 0/4 tested FSWs recognized the E clade version of this epitope PLCFGWCFKL, which differs from the previously defined B clade version by two amino acids, PLTFGWCYKL.
- This epitope was only conserved in CRF01 (subtype E) and subtype B.

**HXB2 Location** Nef (136–145)

Author Location Nef (136–145)

Epitope PLTFGWCYKL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords rate of progression, acute/early infection

References Day et al. 2001

 The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP) • 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.

**HXB2 Location** Nef (136–145)

**Author Location** 

Epitope PLTFGWCYKL

**Epitope name** Nef-PL10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

References Sabbaj et al. 2003

• Among HIV+ individuals who carried HLA A02, 3/29 (10%) recognized this epitope.

**HXB2 Location** Nef (136–145)

**Author Location** Nef (136–145 BRU)

Epitope PLTFGWCYKL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords binding affinity, epitope processing

References Choppin et al. 2001

- Seventy-three 8-11 mer peptides were generated based on putative anchor motifs in the Nef BRU sequence for HLA A2, A3, A24, B7, B8, and B35; these 73 peptides had 81 motifs. 54% bound to the predicted HLA molecule, particularly A2, B7/35, and B8. The strength of HLA-peptide binding was not directly related to the number of individuals that recognized a protein.
- 20s proteasome cleavage of the Nef protein positions 66-100 showed a large fraction of peptides were cleaved ending at: 87L, 83A, 81Y, 71P, 68F and 67G. The frequency of recognition may be in part dictated by the cleavage step in processing.
- PLTFGWCYKL was recognized in 9/28 (32%) of individuals with HLA A2. It was a low affinity HLA-A2 binder.

**HXB2 Location** Nef (136–145)

**Author Location** Nef (136–145)

Epitope PLTFGWCYKL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (A2)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+T-cell epitopes was induced after a fourth boost injection. This was one of the most highly recognized of the 31 peptides that were shown to elicit a response.

**HXB2 Location** Nef (136–145) **Author Location** Nef (136–145)

Epitope PLTFGWCYKL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

 The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized in chronic infection.

**HXB2 Location** Nef (136–145)

**Author Location** Nef (136–145 HXB2)

Epitope PLTFGWCYKL
Subtype B, CRF01\_AE

Immunogen HIV-1 infection Species (MHC) human (A2)

Country Viet Nam.

Assay type HLA binding

**Keywords** subtype comparisons, computational epitope prediction, variant cross-recognition or cross-

neutralization, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-All, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- CRF01\_AE variant plCfgwcFkl had a higher HLA-binding score than the HXB2 epitope.

**HXB2 Location** Nef (136–146)

**Author Location** Nef (136–146 LAI)

**Epitope** PLTFGWCYKLV

Subtype B

**Immunogen** in vitro stimulation or selection

Species (MHC) human (A\*0201)

**Keywords** epitope processing

References Lucchiari-Hartz et al. 2000

- Five naturally processed MHC class I ligands were identified in Nef in the conserved immunogenic region Nef between 123-152.
- All five could be transported by TAP, and 4/5 had N-termini
  that were major cleavage points for the proteasome, only one
  had extended precursor fragments.
- The CTL that recognized PLTFGWCYKL also recognized PLTFGWCYKLV, and both forms of the epitope are naturally processed and both seem to be the direct product of a proteasomal digest, although in low copy number.

**HXB2 Location** Nef (137–145)

Author Location Nef (139–147 HXB3)

Epitope LTFGWCFKL

Immunogen vaccine

Vector/Type: DNA, peptide Strain: B clade HXB3 HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (A\*0201)

 ${\bf Keywords} \ \ {\bf binding} \ {\bf affinity}, computational \ {\bf epitope} \ {\bf predictional}$ 

References Sandberg et al. 2000

- Ten Nef 9-mer peptides were predicted to have a strong binding affinity with HLA-A\*0201 of these, four did bind strongly by a T2 class I stabilization assay, several others bound weakly.
- A CTL immune response to only 3/10 peptides was detected by a 51Cr-release assay after immunization of HLA-A201 transgenic mice with either nef DNA under the control of a CMV promotor, coated on gold particles delivered to abdominal skin by gene gun – LTFGWCFKL did not elicit a CTL response.
- LTFGWCFKL was also tested by subcutaneous injection of Nef peptides in Freund's adjuvant, because it bound strongly to HLA-A\*0201, and the peptide vaccination did elicit a response.
- The lack of response to the nef DNA vaccine and the response to the peptide suggests LTFGWCFKL may not be processed.

HXB2 Location Nef (137–145)

**Author Location** Nef (137–)

Epitope LTFGWCFKL

**Epitope name** Nef137

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Assay type** CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This epitope was one of the previously identified HLA-A2 epitopes studied.
- 3/17 HIV-infected HLA-A2+ people recognized this epitope.

**HXB2 Location** Nef (137–145)

**Author Location** Nef (137–145)

Epitope LTFGWCYKL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (A2)

Assay type proliferation, CD8 T-cell Elispot - IFNγ,

Chromium-release assay

Keywords vaccine-induced epitopes, variant cross-

recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This was one of the most highly recognized of the 31 peptides that were shown to elicit a response.

HXB2 Location Nef (137–145)
Author Location Nef (137–145)
Epitope LTFGWCFKL
Epitope name Nef137-145
Immunogen HIV-1 infection
Species (MHC) human (A2)
Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords responses in children, immunodominance,

characterizing CD8+ T cells

References Chandwani et al. 2004

- Responses to five HLA-A2 presented epitopes in Nef were characterized in a population of 19 HIV infected adults and 21 children. The CD8 T-cell response to Nef was stronger and broader in adults than children, averaging 652 and 87 SFCs/10<sup>6</sup> PBMC, and 2.6 and 0.9 recognized epitopes per person, respectively.
- This epitope was second in an immunodominance hierarchy of the five A02 Nef epitopes studied.

HXB2 Location Nef (137–145)

Author Location Nef (158–166)

Epitope LTFGWCFKL

Immunogen HIV-1 infection

Species (MHC) human (A2 supertype)

Keywords supertype, rate of progression
References Propato et al. 2001

- Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.
- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind all five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802)

**HXB2 Location** Nef (137–145)

**Author Location** Nef

**Epitope** LTFGWCFKL

Epitope name LL9 Subtype B, C

Immunogen HIV-1 infection

**Species (MHC)** human (B\*1517, B57) **Donor MHC** A\*36, A\*66, B\*1517, B\*53, Cw\*04, Cw\*06;

A\*02, A\*23, B\*35, B\*57, Cw\*04, Cw\*07

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** cross-presentation by different HLA, optimal epitope

References Frahm et al. 2005

- HLA-B63-positive subjects were shown to be able to generate CTL responses early in acute HIV infection and to control HIV replication in the absence of antiretroviral treatment. Since HLA-B63 shares the epitope binding motif of HLA-B57 and -B58, it was shown that HLA-B63-positive individuals mounted CTL responses to previously identified B57-restricted epitopes, as well as novel, B63-restricted epitopes. Moreover, these novel B63-restricted epitopes can also be presented by HLA-B57 and -B58
- Optimal epitope was defined in 2 people, 1 carrying HLA-B\*1517(B63), the other carrying B57.

**HXB2 Location** Nef (137–145)

**Author Location** 

**Epitope** LTFGWCFKL

Immunogen

Species (MHC) human (B57)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

• C. Brander notes that this is an B57 epitope.

**HXB2 Location** Nef (137–145)

**Author Location** 

Epitope LTFGWCFKL

Immunogen

Species (MHC) human (B63)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes that this is an B63 epitope.

**HXB2 Location** Nef (137–146)

**Author Location** Nef (137–146)

Epitope LTFGWCYKLV

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6901)

Country India.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, computational epitope prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** Nef (137–146) **Author Location** Nef (221A)

**Epitope** LTFGWCFKLV **Epitope name** Nef-221a

HIV Molecular Immunology 2006/2007

Immunogen HIV-1 infection Species (MHC) human (A2)

**Keywords** binding affinity, subtype comparisons, supertype, computational epitope prediction

References Altfeld et al. 2001c

- HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.
- Three additional previously described HLA-A2 epitopes were added to the set of 20, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- 1/22 individuals with chronic HIV-1 infection recognized this epitope in ELISPOT.
- 2/12 acutely infected individuals recognized this epitope.
- LTFGWCFKLV binds to five HLA-A2 supertype alleles: A\*0203, A\*0201 (highest affinity), A\*0206, A\*6802 and A\*0202.

**HXB2 Location** Nef (137–146)

**Author Location** Nef (137–146)

**Epitope** LTFGWCFKLV

Epitope name LV10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, Flow cytometric T-cell cytokine

assay

**Keywords** optimal epitope **References** Allen *et al.* 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- This epitope did not vary.

**HXB2 Location** Nef (137–146)

**Author Location** Nef (137–146)

Epitope LTFGWCFKLV

Epitope name LV10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Donor MHC** A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Flow cytometric T-cell cytokine

assay

**Keywords** escape, TCR usage, variant cross-recognition or cross-neutralization, characterizing CD8+ T cells

References Allen et al. 2005b

- 4/14 optimal HIV-1 T-cell epitopes in a subject underwent mutation associated with dramatic loss of the original CD8 response. For 1 of the escape variants, a novel CD8 T-cell response equal in magnitude to the wildtype, was generated. CD8 T-cell recognizing the variant epitope utilized a distinct T-cell receptor and did not exhibit any cross-reactivity against the wildtype.
- This epitope did not vary over time.

**HXB2 Location** Nef (137–146)

**Author Location** Nef (137–146)

**Epitope** LTFGWCFKLV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Donor MHC A2, A11, B18, B44, Cw5, Cw12

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** Nef (137–146)

**Author Location** Nef (135–146)

**Epitope** LTFGWCFKLV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A2)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release

assay, HLA binding

Keywords acute/early infection, optimal epitope

References Altfeld et al. 2005

 The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized during chronic infection.

**HXB2 Location** Nef (137–146)

**Author Location** Nef (158–167)

**Epitope** LTFGWCFKLV

Immunogen HIV-1 infection

**Species (MHC)** human (A2 supertype)

Keywords supertype, rate of progression

References Propato et al. 2001

 Long-term nonprogressors (LTNPs) had strong memory resting CD8+ T-cell responses against the majority of epitopes tested, (18 for the A2 supertype, 16 for the A3 supertype) while the effector cells of long-term nonprogressors recognized far fewer epitopes.

- Progressors had memory resting CD8+ T-cells that recognized far fewer epitopes than LTNPs.
- A positive correlation between effector CD8+ T-cells and plasma viremia and a negative correlation between CD8+ effector T-cells and CD4+ T-cells was observed, which may contribute to the inability of LTNPs to clear virus.
- This epitope can bind all five HLA-A2 supertypes alleles (A\*0201, A\*0202, A\*0203, A\*0206 and A\*6802)
- Tetramer staining with A2, beta2microglobulin, and either SLYNTVATL, KLVGKLNWA, or LTFGWCFKL revealed that tetramers detected more HIV-specific sells in LTNP than in progressors, activated effector cells were the minority population, and ELISPOT correlated better with the effector cell subpopulation than the total tetramer stained population.

**HXB2 Location** Nef (141–148)

Author Location Nef (141-)

Epitope WCFKLVPV

**Epitope name** Nef141

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: Nef Adjuvant: Incomplete Freund's Adjuvant (IFA)

**Species (MHC)** human, transgenic mouse (A2)

Assay type CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This peptide was an intermediate A2 binder, and induced a CD8+ T-cell IFN gamma response in 1/6 mice. Responses were detected in 2/17 HIV+ HLA-A2 subjects.

**HXB2 Location** Nef (162–181)

**Author Location** Nef (161–180)

Epitope TSLLHPVSLHGMDDPEREVL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1995

• HIV-specific CTL lines developed by *ex vivo* stimulation with peptide.

**HXB2 Location** Nef (162–181)

**Author Location** Nef (161–180 SF2)

**Epitope** TSLLHPVSLHGMDDPEREVL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- One of these 11 had CTL response to this peptide.

**HXB2 Location** Nef (162–181)

Author Location Nef (101–120 SF2)

Epitope TSLLHPVSLHGMDDPEREVL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997b

 CTL expanded ex vivo were later infused into HIV-1 infected patients.

**HXB2 Location** Nef (162–181)

**Author Location** Nef (161–180 SF2)

Epitope TSLLHPVSLHGMDDPEREVL

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- One of these 11 had CTL response to this peptide.

**HXB2 Location** Nef (166–177)

**Author Location** Nef (160–179 SF2)

Epitope HPVSLHGMDDPE

Immunogen HIV-1 infection

Species (MHC) human (B35)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-B35+ individuals that had a CTL response to this epitope broken down by group: 1/2 group 1, 0/2 group 2, and 0/1 group 3.

HXB2 Location Nef (172–191)

**Author Location** Nef (171–190 SF2)

Epitope GMDDPEREVLEWRFDSRLAF

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-A2, B21.

**HXB2 Location** Nef (175–184)

Author Location Nef (175–184)

Epitope DPEKEVLQWK
Immunogen HIV-1 infection
Species (MHC) human (B7)
References Jin et al. 2000b

- This a B7 epitope, a subdominant CTL response, was defined by an un-conventional approach used to predict epitopes in an HLA B7+ long-term non-progressor.
- Three additional sub-dominant HLA B7 epitopes were defined using EpiMatrix, a non-anchor based strategy for defining potential epitopes, which highlighted 2078 possible epitopes in the autologous HIV-1 derived from the study subject, followed by B7 anchor residue prediction which narrowed the set to 55 peptides, three of which could serve as functional CTL epitopes.

**HXB2 Location** Nef (175–184)

**Author Location** Nef

Epitope DPEKEVLQWK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B7)

**Donor MHC** A\*0301, A\*2301, B\*0702, B\*1503

Country United States.

Keywords escape, acute/early infection

References Bernardin et al. 2005

- Full HIV genomes from nine individuals were analyzed for mutations prior to seroconversion and 7 to 28 days later. It was found that the influence of the host's HLA type was reflected within weeks of infection; a statistically significant number of early nonsynonymous mutations were observed within previously reported CTL epitopes.
- K to E mutation was observed in position 10.

**HXB2 Location** Nef (180–189)

Author Location Nef (180-189 LAI)

Epitope VLEWRFDSRL

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (A\*0201)

References Haas et al. 1996; Haas et al. 1997

- There was a high degree of variation in three CTL epitopes in Nef in four slow and non-progressors, and variant specific CTLs arose over time to eliminate variants, indicating immune selection.
- Noted in Brander et al., 1999 this database, to be A\*0201.

**HXB2 Location** Nef (180–189)

Author Location Nef (180–189 LAI)

Epitope VLEWRFDSRL

Subtype B

Immunogen

Species (MHC) human (A\*0201)

Keywords optimal epitope

References Frahm et al. 2007

• C. Brander notes this is an A\*0201 epitope.

**HXB2 Location** Nef (180–189)

Author Location Nef (180-189 LAI)

Epitope VLMWQFDSRL

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: natural variants HIV component: Nef Adjuvant:

Complete Freund's Adjuvant (CFA)

**Species (MHC)** transgenic mouse (A\*0201)

**Keywords** binding affinity, vaccine-specific epitope characteristics

References Boissonnas et al. 2002

- Ten naturally occurring variants of this epitope were tested for their affinity to HLA-A\*0201 and for their ability to induce gamma-IFN and cytotoxic functions through vaccination of HLA-A\*0201 transgenic mice.
- Only two variants could induce vaccine responses: VLMWQFDSRL, a high affinity binder, and VLQWRFDSRL a medium affinity binder to A\*0201.
- In vivo priming with Nef peptide VLMWQFDSRL induced cross-reactive CTL to 6/7 peptides tested (AlmwKfdsKl, vlmwKfdsrl, vlmwKfdsKl, vlQwRfdsKl, vlVwrfdTrl, and vlAwKLdsrl but not the LAI peptide vlEwrfdsrl)
- In vivo priming with Nef peptide VLQWRFDTRL induced cross-reactive CTL to 3/6 variant Nef peptides (vlMwQfdsrl, vlqwrfdSrl and vlEwrfdsrl).

**HXB2 Location** Nef (180–189)

**Author Location** Nef (190–198)

Epitope VLEWRFDSRL

Immunogen vaccine

Vector/Type: DNA HIV component: HIV-1

Species (MHC) mouse (A\*0201)

**Keywords** epitope processing, vaccine-specific epitope characteristics, immunodominance

References Singh et al. 2002; Sykes & Johnston 1999

- C3H (H-2k) transgenic mice carrying a fused HLA-A\*0201 alpha1 and alpha2 and H-2Dk alpha3 hybrid class I molecule were immunized using an epidermal gene gun with an ubiquitin expression library of 32 plasmids that spanned the HIV-1 genome. Ubiquitin targets the expressed HIV-1 peptides to the proteasome.
- A single immunization with the UB-HIV-1 library vaccine induced potent, stable and multivalent CTL responses against all library members.
- Immunodominant epitopes SLYNTVATL (Gag), ILKEPVHGV (Pol), RIQRGPGRAFVTIGK (Env) and AFHHVAREK (Nef) elicited strong CD8+/IFN- responses and stimulated CTL that were functional in a Cr-release assay and against wild type antigen.
- The presence of multiple plasmids HLA-A\*0201-restricted CTL epitopes did not decrease CTL immunogenicity, and CTL responses to single peptide immunizations were comparable to responses based on mixtures of either 16 or 32 peptides.

**HXB2 Location** Nef (180–189)

Author Location Nef (180–189)

Epitope VLEWRFDSRL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot granzyme B

**Keywords** Th1, characterizing CD8+ T cells **References** Kleen *et al.* 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- Three of seven patients responded to this peptide with GzB producing cells, while one of the three patients responded with IFN-gamma producing cells.

HXB2 Location Nef (180–189)

**Author Location** Nef (180–189)

 ${\bf Epitope} \ \ {\tt VLEWRFDSRL}$ 

Immunogen in vitro stimulation or selection

Species (MHC) human (A2)

Keywords binding affinity, dendritic cells, Th1

References Wilson et al. 1999b

- Dendritic cells are the most potent for priming T cell responses
   DCs can stimulate autologous CTL responses from T cells cultured from HIV negative donors.
- Th1-biasing cytokines IL-12 or IFN alpha enhance CTL responses *in vitro* whether the epitope is delivered by pulsing from peptide, or expressed from within.
- B7 and A2 Nef epitopes were studied and the relative binding affinity of A2 epitopes for A2 was: PLTFGWCYKL greater than VLEWRFDSRL which was much greater than AFHH-VAREL.

**HXB2 Location** Nef (180–189)

**Author Location** Nef (180–189)

**Epitope** VLEWRFDSRL

Immunogen vaccine

Vector/Type: DNA prime with vaccinia boost

Species (MHC) human (A2)

References Woodberry et al. 1999

- A polyepitope vaccine was generated in a vaccinia construct that contiguously encoded seven epitopes, all presented by HLA A-2.
- HHD mice have a transgene of HLA A2 linked to the transmembrane and cytotoxic domains of H-2D<sup>d</sup> – this transgene is the only MHC molecule expressed in the mice.
- CTL responses to Gag (77-85) SLYNTVATL, Pol (476-484) ILKEPVHGV, gp120 (120-128) KLTPLCVTL, and Nef (190-198) AFHHVAREL were observed in HIV polytope HHD-vaccinated mice, and these responses were enhanced with vaccinia boost.
- No CTL immune responses were generated against HLA A2restricted HIV epitopes Nef 157-166 (PLTFGWCYKL), Pol 346-354 (VIYQYMDDL), and Nef 180-189 (VLEWRFDSRL)
- Sixteen HLA A2+ patients were tested for their ability to make CTL responses by peptide restimulation in culture with the epitopes selected for inclusion in the polytope – one individual recognized all seven of these epitopes; 7 patients had CTL cultures able to recognize at least one of the epitopes, and 6 of those 7 recognized more than one epitope, but they were not able to test all peptides for all patients; many patients only had three peptides tested.
- VLEWRFDSRL was recognized by 2 of the HLA-A2 patients.

**HXB2 Location** Nef (180–189)

Author Location Nef (180–189 LAI)

Epitope VLEWRFDSRL

Epitope name N3

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (A2)

**Keywords** HAART, ART

References Mollet et al. 2000

- A panel of 16 epitopes covering 15 class I alleles was tested in 14 HIV+ patients from an unselected Caucasian population treated with HAART, using CD8+ cell IFNgamma production to measure responses.
- In general, during the first month of treatment viral load decreased and frequencies of HIV-specific CTL tripled and broadened eight new HIV specificities that were not previously detectable were newly detected, as were CMV specific CD8+PBL but with continued viral suppression, HIV-specific responses diminished.
- Viral rebounds gave different patterns of response: increases or decreases in pre-existing response, new specificities, or no change.

**HXB2 Location** Nef (180–189)

**Author Location** Nef (179–188 93TH253 subtype CRF01)

Epitope VLEWRFDSRL

Subtype CRF01 AE

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords subtype comparisons

References Bond et al. 2001

- More than half of a cohort of HIV+ female sex workers (FSW) from Northern Thailand were HLA-A11 positive, and this study concentrated on A11 epitopes in this group, although E clade versions of previously defined B-clade A2 and A24 epitopes were also tested.
- 0/4 tested FSWs recognized the E clade version of this epitope VLIWKFDSAL, which differs from the previously defined B clade version by three amino acids, VLEWRFDSRL.

HXB2 Location Nef (180–189)

**Author Location** Nef (180–189)

Epitope VLEWRFDSRL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords rate of progression, acute/early infection

References Day et al. 2001

- The CTL response to optimally defined CTL epitopes restricted by HLA class I A and B alleles in individuals who coexpressed HLA A2, A3, and B7 was studied in eight HIV-1-infected subjects, two with acute infection, five with chronic, and one long-term non-progressor (LTNP)
- 2 to 17 epitopes were recognized in a given individual, A2-restricted CTL response tended to be narrow and never dominated the response, and 25/27 epitopes were targeted by at least one person.

**HXB2 Location** Nef (180–189)

Author Location Nef (178–187)

Epitope VLEWRFDSRL

Immunogen HIV-1 infection Species (MHC) human (A2) Country Spain.

**Assay type** proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow

cytometric T-cell cytokine assay

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- 5/19 patients recognized this epitope.

HXB2 Location Nef (180–189) Author Location Nef (180–189) Epitope VLEWRFDSRL Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2) Country United States.

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, HLA binding

**Keywords** acute/early infection, optimal epitope

References Altfeld et al. 2005

 The most frequently targeted HLA-A2-restricted CD8+ T-cell epitopes in chronic infection were significantly less frequently recognized during primary infection. This epitope was only recognized in chronic infection, and even then was recognized infrequently.

HXB2 Location Nef (180–189) Author Location Nef (180–189 HXB2)

Epitope VLEWRFDSRL
Subtype B, CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A2)
Country Viet Nam

Country Viet Nam.
Assay type HLA binding

**Keywords** subtype comparisons, computational epitope prediction, variant cross-recognition or cross-

neutralization, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-All, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- CRF01\_AE variant lMwKfdsAl had a higher HLA-binding score than the HXB2 epitope, which isn't predicted to bind to A2.

HXB2 Location Nef (181–189) Author Location Nef (181–189) Epitope LEWRFDSRL Epitope name Nef181-189 Immunogen HIV-1 infection Species (MHC) human (A2) Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** responses in children, immunodominance, characterizing CD8+ T cells

References Chandwani et al. 2004

- Responses to five HLA-A2 presented epitopes in Nef were characterized in a population of 19 HIV infected adults and 21 children. The CD8 T-cell response to Nef was stronger and broader in adults than children, averaging 652 and 87 SFCs/10<sup>6</sup> PBMC, and 2.6 and 0.9 recognized epitopes per person, respectively.
- This was not the immunodominant response.

HXB2 Location Nef (182–189) Author Location Nef (182–189) Epitope EWRFDSRL Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (B8)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes, variant cross-recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of the 31 epitopes that the vaccinated volunteers responded to.

HXB2 Location Nef (182–198)
Author Location Nef (182–198 BRU)
Epitope EWRFDSRLAFHHVAREL

Immunogen HIV-1 infection Species (MHC) human (A1, B8)

**References** Hadida *et al.* 1992

 HIV-1 specific CTLs detected in lymphoid organs of HIV-1 infected patients.

HXB2 Location Nef (182–198)

Author Location Nef (182–198 LAI)

Epitope EWRFDSRLAFHHVAREL

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2, A25) References Hadida *et al.* 1995

 The C-terminal region of Nef (182-205) contains multiple CTL epitopes with 5 distinct HLA restrictions.

**HXB2 Location** Nef (182–198) **Author Location** Nef (182–198 BRU) Epitope EWRFDSRLAFHHVAREL Immunogen HIV-1 infection Species (MHC) human (A25) References Cheynier *et al.* 1992

• CTL isolated in children born to HIV-1 positive mothers.

HXB2 Location Nef (182–198)

Author Location Nef (182–198 LAI)

Epitope EWRFDSRLAFHHVAREL

Subtype B

Immunogen HIV-1 infection

Immunogen HIV-1 infection Species (MHC) human (B35) References Hadida *et al.* 1995

 The C-terminal region of Nef (182-205) contains multiple CTL epitopes with 5 distinct HLA restrictions.

HXB2 Location Nef (182–198)

Author Location Nef (182–198 LAI)

Epitope EWRFDSRLAFHHVAREL
Subtype B

Immunogen vaccine

Vector/Type: vaccinia, Mengo virus Strain: • B clade LAI HIV component: Nef

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Van der Ryst et al. 1998

- Macaca mulatta did not have a detectable response to Rec Mengo virus-HIV-1 Nef 65-206 vaccine.
- BALB/c mice had a weak response to this epitope in the Mengo virus construct – in contrast, HIV-1 Nef induces a strong CTL response in mice when presented in a vaccinia background.

HXB2 Location Nef (182–201) Author Location Nef (191–205 SF2) Epitope EWRFDSRLAFHHVARELHPE

Immunogen HIV-1 infection

Species (MHC) human

References Lieberman et al. 1997a

- Of 25 patients, most had CTL specific for more than 1 HIV-1 protein.
- Eleven subjects had CTL that could recognize vacciniaexpressed LAI Nef.
- One of these 11 had CTL response to this peptide.
- The responding subject was HLA-A2, B21.

HXB2 Location Nef (182–205) Author Location Nef (182–205 LAI)

**Epitope** EWRFDSRLAFHHVARELHPEYFKN

Subtype B Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long amino acid peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I trial
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees – 4/10 reacted to this Nef peptide.

- 9/12 tested mounted a CTL response to at least one of the six peptides; each of the six peptides elicited a CTL response in at least one individual.
- None of the 12 tested had an IgG response to this peptide.

**HXB2 Location** Nef (183–191)

**Author Location** 

Epitope WRFDSRLAF
Epitope name Nef-WF9
Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*1503)

**Donor MHC** A\*2904 A\*3002 B\*1503 B\*5802 Cw\*0202 Cw\*0602

**Keywords** HAART, ART **References** Sabbaj *et al.* 2003

- This study monitored epitope responses in HIV-1 infected minority women living in the United States.
- 24 epitopes were described 8 were novel, 8 used new restricting elements but were previously defined epitopes, and 8 were previously described.
- Serial peptide truncations were used to define optimal epitopes for CTL cell lines isolated from 12 individuals, assayed by a Cr-release.
- This epitope was newly defined in this study.
- Subject 01RCH50 also recognized the epitope RMRGAHT-NDV, RT(356-365), A\*3002 – she was African American, was on HAART, had a viral load of 960 and CD4 count of 728.
- Among HIV+ individuals who carried HLA B15, 3/17 (18%) recognized this epitope.

**HXB2 Location** Nef (183–191)

**Author Location** Nef (183–191)

Epitope WRFDSRLAF

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*1503)

**Keywords** optimal epitope **References** Frahm *et al.* 2007

**HXB2 Location** Nef (183–191)

**Author Location** Nef (183–191)

Epitope WRFDSRLAF

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*1503)

**Donor MHC** A\*2301, B\*3501, B\*1503, Cw2, Cw7

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** binding affinity, acute/early infection, early-expressed proteins

References Cao et al. 2003

• All proteins were scanned and optimal epitopes were mapped in a study of CD8+ gamma IFN T-cell responses in 21 men within 15-92 days post-HIV-1 infection. Subjects initially a showed narrow immune responses with a mean of 2.3 epitopes recognized per patient. The immune response broadened later in infection. No correlation between the plasma viral load and the number of recognized epitopes or the frequency of IFN-gamma secreting cells was observed, and there was no correlation between the functional avidity of a responses and the abundance of IFN-secreting cells that recognized the epitope. The first epitopes to be recognized did not tend to be the most avid, and earliest responses tended to be directed against Nef, Tat, Vpr, and Env.

- All HIV-1 proteins except Vpu were recognized, and a responses to a total of 41 optimal epitopes were characterized;
   24 had been identified previously. 48% of the total T-cell responses were directed against Nef and 43% were directed against Gag.
- More common HLA alleles were less frequently used in primary infection than less common alleles, for example A3, B35, B57, and B62 were more frequently utilized than A1, A2, A30, and A44.

**HXB2 Location** Nef (183–191)

**Author Location Nef** 

**Epitope** WRFDSRLAF

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (B\*1503)

**Donor MHC** A\*0301, A\*2301, B\*0702, B\*1503

Country United States.

Keywords escape, acute/early infection

References Bernardin et al. 2005

- Full HIV genomes from nine individuals were analyzed for mutations prior to seroconversion and 7 to 28 days later. It was found that the influence of the host's HLA type was reflected within weeks of infection; a statistically significant number of early nonsynonymous mutations were observed within previously reported CTL epitopes.
- K to E mutation was observed in position 2.

**HXB2 Location** Nef (183–192)

**Author Location** Nef (183–192)

**Epitope** WRFDSRLAFH

Subtype B

Immunogen HIV-1 infection Species (MHC) human (A1)

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- This epitope was reactive, but escape mutations did not accrue in it over time.

**HXB2 Location** Nef (184–191)

Author Location Nef (184–191 HXB2)

Epitope RFDSRLAF

Subtype B, CRF01\_AE

Immunogen HIV-1 infection

Species (MHC) human (A1)

Country Viet Nam.

Assay type HLA binding

Keywords subtype comparisons, computational epitope prediction, variant cross-recognition or cross-neutralization, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-A11, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- CRF01\_AE common variant KfdsAlaR had same HLA-binding score as the HXB2 epitope.

**HXB2 Location** Nef (186–193)

Author Location Nef (186–193 LAI)

**Epitope** DSRLAFHH

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35)

References Hadida et al. 1995

 The C-terminal region of Nef (182-205) contains multiple CTL epitopes with 5 distinct HLA restrictions.

**HXB2 Location** Nef (186–194)

**Author Location** Nef (186–194)

**Epitope** DSRLAFHHM

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A24)

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2001a

 ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** Nef (186–194)

Author Location Nef (186–194)

Epitope DSRLAFHHM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A24)

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot granzyme B

**Keywords** Th1, characterizing CD8+ T cells

References Kleen et al. 2004

- Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.
- Two of seven patients responded to this peptide with GzB producing cells, while none of the patients responded with IFN-gamma producing cells.

**HXB2 Location** Nef (186–194)

Author Location Nef (186–194)

**Epitope** DSRLAFHHM **Epitope name** DM9

Subtype B

**Immunogen** HIV-1 infection **Species (MHC)** human (A24)

**Donor MHC** A\*24, A\*31, B\*47, B\*15, Cw\*04, Cw\*07; A\*24, A\*30, B\*39, B\*47, Cw\*12, Cw\*17; A\*24, A\*23, B\*39, B\*07, Cw\*12, Cw\*17

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant transmission, escape, acute/early infection, rever-

sion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs. Escape variants were commonly detected in maternal plasma. Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses
- DSRLAFHHM is the known A24 epitope. A known escape variant, DSRLAFqHM, was transmitted from an A24- mother to her A24+ infant, where it was gradually lost over time, present in 6/10 clones at 2 months, 7/10 at 4 months, and 0/10 at 15 months.
- Another escape variant was present in an A24+ mother, DSt-LAFqHk and this form was transmitted to her A24+ infant where it persisted in 30/30 sequences sampled over 12 months.
- DSRLAFHHM had higher responder cell frequencies in the A24+ mother than DStLAFqHk. Her A24+ infant did not recognize either form. The variant DSRLAFqHM also stimulated lower responder cell frequencies.

**HXB2 Location** Nef (186–194)

Author Location Nef (186–194 BRU)

Epitope DSRLAFHHV

Immunogen

Species (MHC) human (B51)

References Connan et al. 1994

• Resulted in the assembly of HLA-B51.

**HXB2 Location** Nef (186–194)

**Author Location** Nef (186–194)

**Epitope** DSRLAFHHV

Immunogen HIV-1 infection

Species (MHC) human (B51)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4, •

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 7 epitopes from this individual that varied over time, although the internal mutations did not become fixed. The dsLlaLRhM variant residues arose at early time points, and the dsrlaVhhv variant residue arose at intermediate time points.

**HXB2 Location** Nef (186–194)

**Author Location Nef** 

**Epitope** DSRLAFHHV

Epitope name DV9

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A28, A29, B14, B44, Cw8

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** subtype comparisons, escape, characterizing CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- An escape mutation at position 7, DSRLAFqHV, was found not to correspond to the most polymorphic residue in the epitope. This is a novel unmapped epitope.

**HXB2 Location** Nef (188–196)

**Author Location** Nef (192–200 SF2)

**Epitope** KLAFHHMAR

Subtype B

**Immunogen** HIV-1 infection, computer prediction

Species (MHC) human (A\*3303)

Assay type Chromium-release assay

**Keywords** binding affinity, computational epitope prediction

References Hossain et al. 2003

- HLA-A\*3303 is a common HLA allele in east and southeast Asia. Pol, Gag and Nef SF2 proteins were scanned for potential A\*3303 epitopes. 99 potential epitopes were synthesized, and 52/99 bound to A\*3303. Six of these served as peptide-targets for lysis by PBMC from infected individuals, and clones derived from 4 of these 6 could lyse HIV-vaccinia infected target cells, indicating proper processing.
- This epitope is one of the 2/6 peptides that could induce CTL responses in the PBMC of infected individuals, but was not properly processed in a vaccinia-HIV infected target cell.

**HXB2 Location** Nef (188–196)

Author Location Nef (188–196 LAI)

**Epitope** RLAFHHVAR

Subtype B

Immunogen HIV-1 infection Species (MHC) human (B52)

References Hadida et al. 1995

 The C-terminal region of Nef (182-205) contains multiple CTL epitopes with 5 distinct HLA restrictions.

HXB2 Location Nef (188–201) Author Location Nef (188–201 LAI)

Epitope RLAFHHVARELHPE

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (B35, Cw4)

References Buseyne et al. 1993a

- Vertical transmission of HIV ranges from 13% to 39%
- Primary assays showed that cytotoxic activity against at least one HIV protein was detected in 70% of infected children.
- Epitopes recognized in five children were mapped using synthetic peptides and secondary cultures.
- Patient EM13, who had a CTL response to three epitopes in Nef, was infected via blood transfusion after birth and went from CDC stage P2A to P2E during the study.

**HXB2 Location** Nef (189–198)

Author Location Nef (189–198)

**Epitope** LAFHHVAREL

Subtype C

Immunogen HIV-1 infection

Species (MHC) human (A\*6901)

Country India.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords subtype comparisons, computational epitope

prediction, immunodominance

References Thakar et al. 2005

• PBMCs from HIV-1 infected Indian subjects were tested for HIV-1 Gag (with C clade peptides), Nef (with B clade peptides), and Env (clade unspecified) specific T-cell responses. 5 immunodominant regions conserved across clades were identified in Gag and Nef. 3 antigenic regions were found to be recognized by CTLs in the Indian subjects; these regions were not identified as immunodominant in studies from Africa. 26 epitopes were predicted within reactive peptides, of which 90% were clustered in the conserved immunodominant regions of Gag and Nef. 3 of the epitopes were highly conserved across clades, and 7 novel epitopes were found.

**HXB2 Location** Nef (190–198)

**Author Location** Nef

Epitope AFHHVAREL

Epitope name Nef AL9

Immunogen HIV-1 infection

Species (MHC) human (A\*0201)

Keywords subtype comparisons, supertype, computa-

tional epitope prediction

References Altfeld et al. 2001c

HIV was scanned for all peptides which carried the A2-supermotif pattern conserved in more than 50% of B clade sequences – 233 peptides met this criteria, and 30 of these bound to HLA-A\*0201 – 20/30 bound to at least 3/5 of HLA-A2 supertype alleles tested.

- Three additional previously described HLA-A2 epitopes were added to the set of 20, including Nef AL9, and 18/22 chronically infected HLA-A2 individuals had CTL that recognized at least one of the 23 peptides (median of 2 and maximum of 6), while 6/12 acute infected individuals recognized at least 1 (median of 1 and maximum of 2)
- RT VL9 was not recognized by any of the 22 HLA-A2 patients with chronic HIV-1 infection or the 13 HLA-A2 patients with acute HIV-1 infection included in this study.

**HXB2 Location** Nef (190–198)

**Author Location Nef** 

Epitope ALKHRAYEL

Subtype A

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: A clade

HIV component: p17 Gag, p24 Gag

**Species (MHC)** human, macaque (A\*0201)

**Keywords** subtype comparisons, epitope processing, immunodominance

References Hanke & McMichael 2000; Wee et al. 2002

- The HIV-1 subtype A focused vaccine HIVA contains p24 and p17, in a reversed order relative to the Gag polyprotein to prevent myristylation of p17, which could direct the protein to the cell membrane and inhibit efficient peptide processing and class I presentation, as well as a polyepitope string of conserved, often immunodominant epitopes that were selected to have particularly good cross-reactive potential for the A-clade epidemic in Nairobi, Kenya. A DNA and MVA prime-boost vaccination protocol using the HIVA antigen will be used in a phase III clinical trial in Kenya. This epitope is included in the polyepitope string Hanke & McMichael [2000].
- Multiple CD4+ or CD8+ T-cell vaccine-induced responses to peptide pools were detected using intracellular cytokine staining and IFNgamma Elispot assays after vaccination of 5 macaques. The response to the Mamu A\*01 SIV p27 epitope p11C (CTPYDINQM), included in the polyepitope region, was not immunodominant in the Mamu A\*01 vaccinated macaques, possibly because of processing limitations in context of the artificial polyepitope string Wee et al. [2002].

**HXB2 Location** Nef (190–198)

Author Location Nef (190-198 LAI)

Epitope AFHHVAREL

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2)

Keywords subtype comparisons, HIV exposed persis-

tently seronegative (HEPS)

References Rowland-Jones et al. 1998a

- CTL recognition reported in the context of HLA-B52 and A2.1, A2.2 and A2.4.
- A CTL response was found in exposed but uninfected prostitutes from Nairobi using previously-defined B clade epitopes that tended to be conserved in A and D clades such cross-reactivity could protect against both A and D and confer protection in Nairobi where both subtypes are circulating.
- The A subtype consensus is ALKHRAYEL.
- The D subtype consensus is AfEHKAREm.

- Hunziker *et al.* [1998] suggests that HLA-A2 does not in fact present this epitope, and notes that it does not promote A2 assembly Connan *et al.* [1994] also see Brander *et al.* [1998b]
- Hunziker *et al.* [1998] maintains that HLA-A2 does not present this epitope contrary to an earlier report Hadida *et al.* [1995], (also see Brander *et al.* [1998a])—despite the position of Hunziker *et al.*, Rowland-Jones and colleagues are confident that this epitope in its A clade form is presented by HLA-A\*0201 and A\*0202, and it is one of the most common responses seen in both seropositive and exposed-uninfected donors from Nairobi (Rupert Kaul, pers. comm.)

HXB2 Location Nef (190–198) Author Location Nef (190–198)

Epitope AFHHVAREL

Immunogen in vitro stimulation or selection

Species (MHC) human (A2)

Keywords binding affinity, dendritic cells, Th1

References Wilson et al. 1999b

- Dendritic cells are the most potent for priming T cell responses
   DCs can stimulate autologous CTL responses from T cells cultured from HIV negative donors.
- Th1-biasing cytokines IL-12 or IFN alpha enhance CTL responses *in vitro* whether the epitope is delivered by pulsing from peptide, or expressed from within.
- B7 and A2 Nef epitopes were studied and the relative binding affinity of A2 epitopes for A2 was: PLTFGWCYKL greater than VLEWRFDSRL which was much greater than AFHH-VAREL.

**HXB2 Location** Nef (190–198)

**Author Location** Nef (190–198)

**Epitope** AFHHVAREL

Immunogen vaccine

Vector/Type: vaccinia

Species (MHC) human (A2)

References Woodberry et al. 1999

- A polyepitope vaccine was generated in a vaccinia construct that contiguously encoded seven epitopes, all presented by HLA A-2.
- HHD mice have a transgene of HLA A2 linked to the transmembrane and cytotoxic domains of H-2D<sup>d</sup> – this transgene is the only MHC molecule expressed in the mice.
- CTL responses to Gag (77-85) SLYNTVATL, Pol (476-484) ILKEPVHGV, gp120 (120-128) KLTPLCVTL, and Nef (190-198) AFHHVAREL were observed in HIV polytope HHDvaccinated mice, and these responses were enhanced with vaccinia boost.
- No CTL immune responses were generated against HLA A2restricted HIV epitopes Nef 157-166 (PLTFGWCYKL), Pol 346-354 (VIYQYMDDL), and Nef 180-189 (VLEWRFDSRL)
- Sixteen HLA A2+ patients were tested for their ability to make CTL responses by peptide restimulation in culture with the epitopes selected for inclusion in the polytope – one individual recognized all seven of these epitopes; 7 patients had CTL cultures able to recognize at least one of the epitopes, and 6 of those 7 recognized more than one epitope, but they were not able to test all peptides for all patients; many patients only had three peptides tested.
- AFHHVAREL was recognized by 2 of the patients.

**HXB2 Location** Nef (190–198)

**Author Location** Nef (190–198 SF2)

Epitope AFHHVAREL

Immunogen HIV-1 infection

Species (MHC) human (A2)

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the response was determined using ELISPOT by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Previously described and newly defined optimal epitopes were tested for CTL response.
- Number of HLA-A2+ individuals that had a CTL response to this epitope broken down by group: 0/10 group 1, 1/6 group 2, and 0/4 group 3.

**HXB2 Location** Nef (190–198)

Author Location Nef (190-198)

Epitope ALKHRAYEL

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human (A2)

**Keywords** subtype comparisons, HIV exposed persistently seronegative (HEPS)

References Kaul et al. 2001a

- Variants ALKHRAYEL and AFHHVAREL are A/B clade specific.
- ELISPOT was used to study CTL responses to a panel of 54 predefined HIV-1 epitopes in 91 HIV-1-exposed, persistently seronegative (HEPS) and 87 HIV-1-infected female Nairobi sex workers.

**HXB2 Location** Nef (190–198)

Author Location Nef (190-)

Epitope AFHHVAREL

Epitope name Nef190

Immunogen HIV-1 infection

Species (MHC) human (A2)

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** binding affinity, subtype comparisons, computational epitope prediction

References Corbet et al. 2003

- HLA-A2-restricted HIV-1 CTL epitopes were computationally predicted. Binding affinities for HLA-A\*0204, immunogenicity in HLA-A\*0201 transgenic mice, and responses to the peptides in 17 HIV-1 infected patients were tested. 31 novel conserved A2 epitopes were detected. An average of 4 epitopes were recognized per patient.
- This epitope was one of the previously identified HLA-A2 epitopes studied.
- None of the 17 HIV-infected HLA-A2+ people in this study recognized this epitope.

HXB2 Location Nef (190–198)

Author Location Nef (190–198)

Epitope ALHHVAREL
Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade
LAI HIV component: Env, Gag, Nef Adjuvant: QS21

Species (MHC) human (A2)

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Chromium-release assay

**Keywords** vaccine-induced epitopes **References** Gahéry-Ségard *et al.* 2003

After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of 31 that the vaccinated volunteers responded to.

HXB2 Location Nef (190–198)
Author Location Nef (subtype B)
Epitope AFHHVAREL
Subtype B
Immunogen HIV-1 exposed seronegative

Species (MHC) human (A2, A\*0202, A\*0201)

**Keywords** subtype comparisons **References** Rowland-Jones *et al.* 1998b

- HIV-specific CTL were found in exposed seronegative prostitutes from Nairobi these CTL may confer protection.
- Seroprevalence in this cohort is 90-95% and their HIV-1 exposure is among the highest in the world.
- Most isolated HIV strains are clade A in Nairobi, although clades C and D are also found – B clade epitopes are often crossreactive, however stronger responses are frequently observed using A or D clade versions of epitopes.
- Clade A version of the epitope: ALKHRAYEL, Clade D epitope: AFEHKAREM.
- This epitope was recognized by two different exposed and uninfected prostitutes.

HXB2 Location Nef (190–198) Author Location Nef (190–198) Epitope AFHHVAREL Subtype B

Immunogen HIV-1 infection Species (MHC) human (A2, B52) Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Elispot granzyme B

Keywords Th1, characterizing CD8+ T cells

References Kleen et al. 2004

 Only 20% of CD8+ T-cells produce IFN-gamma and granzyme B simultaneously (Tc1a). Two additional subpopulations of HIV specific CD8 cells are found, each one constituting 30-40% of the CD8 cell pool. One of these (Tc1b) secretes IFN-gamma only, and the other one (Tc1c) secretes GzB only.  Three of seven patients responded to this peptide with GzB producing cells, while one of three patients also responded with IFN-gamma producing cells.

HXB2 Location Nef (190–198)
Author Location Nef (190–198 HXB2)
Epitope AFHHVAREL
Subtype B, CRF01\_AE
Immunogen HIV-1 infection
Species (MHC) human (A24)
Country Viet Nam.
Assay type HLA binding

**Keywords** subtype comparisons, computational epitope prediction, escape, variant cross-recognition or cross-neutralization, vaccine antigen design

References Lazaro et al. 2005

- The most common HLA-alleles in the Vietnamese population were found to be HLA-All, A02, A33, B75, B46, and B62. Several epitopes cluster in short regions in Gag and Nef that are presented by these HLA molecules; these epitopes may be useful targets for vaccine antigens. In a comparison of CRF01\_AE sequences to HXB2, 14 mutations were found in epitopes in these Gag and Nef pluriepitopic regions; 7 did not affect the HLA binding score, 3 increased it, and 4 reduced it.
- ArrHiAREL, ArtHiAREL, and ArkHiAREL variants are the three forms found in CRF01, and none are predicted to bind to A24.

HXB2 Location Nef (190–198) Author Location Nef (190–198 LAI) Epitope AFHHVAREK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (A3)

References Hadida et al. 1995

 Naturally occurring L to K anchor substitution abrogates A2 binding, but permits HLA-A3 binding.

HXB2 Location Nef (190–198)
Author Location Nef (190–198)
Epitope AFHHVAREK
Immunogen HIV-1 infection
Species (MHC) human (A3)
Country Spain.

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric T-cell cytokine assay

**Keywords** HAART, ART, supervised treatment interruptions (STI), immune dysfunction

References Plana et al. 2004

- Structured treatment interruption (STI) alone is not able to control viral replication in chronically infected patients, probably due to lack of strong, maintained T-helper cell responses. HIV-1 specific CD8+ T-cell responses were shown to increase significantly until the end of the follow up, but were not correlated with viral load.
- Less than 2 of 14 patients recognized this epitope.

HXB2 Location Nef (190–198) Author Location Nef (190–198) Epitope AFHHVAREK

Epitope name AL9

Immunogen HIV-1 infection Species (MHC) human (A3)

**Donor MHC** A\*03, A\*31, B\*08, B\*15, Cw\*04, Cw\*07

Country United Kingdom.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** responses in children, mother-to-infant trans-

mission, escape, characterizing CD8+ T cells, reversion, viral fitness

References Sanchez-Merino et al. 2005

- CD8 T-cell responses were examined in mother-infant pairs. Escape variants were commonly detected in maternal plasma. Early infant plasma viruses showed heterogeneity of gag and nef gene sequences as well as mother-to-child transmission of CD8 T-cell escape variants. The stability of escape mutants in the infant over time was determined by infant HLA haplotype and viral fitness. Sequential plasma specimens from infants showed changes in CD8 T-cell epitope sequences, suggesting that infants are capable of generating virus-specific CD8 T-cell responses.
- AFHHVAREK is an A3 epitope, and a mixture of variants was present in the mother and in her A3- (A31+) infant at 2 months. One of the variants was AFqHmAREl, and this form was not found in 10 clones at the 15 month time point.

**HXB2 Location** Nef (190–198)

Author Location Nef (190-198)

Epitope AFHHVAREK

Immunogen HIV-1 infection

Species (MHC) human (B51)

Donor MHC A03, A32, B51, B15, Cw03, Cw06, DR4,

DR8, DQ7

Country Netherlands.

Assay type CD8 T-cell Elispot - IFNγ, Flow cytometric

T-cell cytokine assay

Keywords rate of progression, escape

References Geels et al. 2003

- Both stable potential epitopes and epitopes with escape mutations were found in a patient with increasing viral load over a time period of 4 years. Fixation of escape mutations occured sequentially and correlated with the rise in viral load and with the increased number of productively infected cells.
- This is one of 7 epitopes from this individual that varied over time, although the internal mutations did not become fixed. The aLRhMarek variant residues arose at early time points, the aVhhvarek variant residue arose at intermediate time points, and afhhvaXeI variant residues arose at late time points.

**HXB2 Location** Nef (190–198)

**Author Location** 

Epitope ALKHRAYEL Immunogen HIV-1 infection

Species (MHC) human

Keywords HIV exposed persistently seronegative

(HEPS)

References Kaul et al. 2001c

- This study examines CTL responses in HIV exposed, persistently seronegative individuals, HEPS, who eventually seroconverted - 11/114 HEPS Nairobi sex workers eventually seroconverted, and for six of these HIV CTL reactive epitopes had been defined while seronegative.
- The epidemiological factor associated with seroconversion was stopping sex work. HIV-specific CTL activity declined when HEPS sex workers stopped working for a period or retired.
- This epitope was in 1/22 HEPS controls, ML1749.

**HXB2 Location** Nef (190–206)

**Author Location** Nef

Epitope AFRHMARELHPEYYKNC

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A1, A3, B7, B57, Cw6, Cw7

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

Keywords subtype comparisons, escape, characterizing

CD8+ T cells, reversion, viral fitness

References Allen et al. 2005a

- · Two-thirds of all mutations arising in nonenvelope proteins during the transition from acute to chronic infection were found to be due to either CD8 T-cell-associated selective pressures or reversion of HLA-I-associated mutations selected in a previous host. The predominance of escape mutations at highly polymorphic sites was observed across different HIV-1 clades.
- Novel unmapped epitope.
- AFRHMARELHPEYYKNC acquired a substitution over time, AFRHMAREmHPEYYKNC.

**HXB2 Location** Nef (192–206)

Author Location Nef (192–206 BRU)

Epitope HHVARELHPEYFKNC

Immunogen HIV-1 infection

**Species (MHC)** human (A1)

References Hadida et al. 1992

• HIV-1 specific CTLs detected in lymphoid organs of HIV-1 infected patients.

**HXB2 Location** Nef (195–202)

Author Location Nef (195–202)

**Epitope** ARELHPEY Subtype B

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Env, Gag, Nef Ad-

juvant: QS21

Species (MHC) human (A1)

Assay type proliferation, CD8 T-cell Elispot - IFNγ,

Chromium-release assay

Keywords vaccine-induced epitopes, variant cross-

recognition or cross-neutralization

References Gahéry-Ségard et al. 2003

• After injection of an anti-HIV-1 lipopeptide vaccine and the booster injections, a response was observed in >85% of 28 volunteers, and a long-term immune response was observed in >50% of the volunteers. A positive effect of QS21 adjuvant was found for B-and CD4+T-cell responses. A higher number of CD8+ T-cell epitopes was induced after a fourth boost injection. This epitope was one of 31 that were recognized in the vaccinees.

HXB2 Location Nef (195-202) Author Location Nef (195–202 BRU) **Epitope** ARELHPEY

Subtype B, CRF02\_AG Immunogen HIV-1 infection Species (MHC) human (A1) Country Cote D'Ivoire.

> Assay type CD8 T-cell Elispot - IFNγ **Keywords** subtype comparisons

References Inwoley et al. 2005

- CD8+ T-cells from HIV-1 CRF02\_AG-infected Ivorians could recognize clade B epitopes. No difference was observed in the number of recognized peptide pools between nine French subtype B infected study subjects, and nine CRF02\_AG infected Ivorian subjects.
- This epitope was recognized by 0/9 CRF02\_AG-infected patients, and by 1/9 B-infected patients. Sequence variants with two amino acid substitutions were found in 5/6 Ivorian subjects.
- An epitope variant was found in 1/4 French patients, and it happened to be the one patient that recognized the epitope: AREmHPEY.

**HXB2 Location** Nef **Author Location Nef Epitope** Immunogen HIV-1 infection

**Species (MHC)** human (A\*0201, Cw\*08)

References Shacklett et al. 2000

• HIV-1 specific, MHC class I-restricted CTL killing was detected in duodenal and rectal gut associated lymphoid tissue (GALT) sites from three infected individuals – the distribution of class I restricted CTL was different in the peripheral blood samples and GALT samples.

**HXB2 Location** Nef **Author Location Nef Epitope** Subtype B

Immunogen HIV-1 infection Species (MHC) human (B\*35) **Keywords** rate of progression

References Jin et al. 2002

- Patients with HLA-B\*35 variants B\*3502, B\*3503, B\*3504, and B\*5301 tend to proceed to AIDS more quickly than those with B\*3501.
- Of 32 patients with HLA-B\*35 alleles CD8+ CTL responses were quantified using an intracellular cytokine staining assay -75% had responses to Pol, 69% to Gag, 50% to Nef, and 41%to Env.

• The overall magnitude of CTL responses did not differ between those bearing B\*3501 and the others. A higher percentage of Gag responses was observed in those that had lower RNA levels that carried B\*3501, and there was a negative association with viral load and CTL activity. The data is consistent with higher levels of CTL responses contributing to protection in B\*3501 individuals, but not in B\*3502, B\*3503, B\*3504, and B\*5301 individuals.

**HXB2 Location** Nef **Author Location** Nef **Epitope** Subtype B Immunogen vaccine

> Vector/Type: DNA Strain: B clade NL43 HIV component: Nef Adjuvant: Bupivacaine

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type CD8 T-cell Elispot - IFNγ

Keywords class I down-regulation by Nef, vaccine antigen design

References Majumder et al. 2003

- · Non-functional Nef vaccine constructs that do not downregulate class I or CD4 proteins are shown to be capable of inducing primary and memory T cell immune response after DNA vaccination in BALB/c mice, which makes them good candidates for vaccines.
- The responses to peptide pools suggest the C-terminal region of Nef is more immunogenic (the two most reactive peptide pools spanned positions 126-175, and positions 166-215).

**HXB2 Location** Nef **Author Location** Nef (BRU) **Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade BRU HIV component: Nef

**Species (MHC)** mouse (H-2D<sup>d</sup>) References Collings et al. 1999

- A comparison of DNA vaccination with HIV-1 Nef expression vectors pBN-CMV-NEF and pBN-RSV-NEF (self-replicating), pCGE2-NEF (non-replicating).
- CTL immune responses were detected using all three expression vectors, while a humoral immune response to Nef was only observed in the self-replicating expression vectors; possibly antibody responses require higher levels of protein expression.

**HXB2 Location** Nef Author Location Nef (SIV)

**Epitope** 

Immunogen SIV infection

Species (MHC) macaque (Mamu-A\*11, Mamu-B\*03, Mamu-B\*04, Mamu-B\*17)

References Dzuris et al. 2000

Cell binding assays for Mamu molecules were employed to describe the peptide binding motifs for Mamu-A\*11, -B\*03, -B\*03, -B\*04, and -B\*17 CTL epitopes - a similarity for Mamu-A\*11 and -B\*03 and human HLA-B\*44 and -B\*27, respectively, was observed – all epitopes studied were SIV epitopes, so not specifically listed here.

**HXB2 Location** Nef

Author Location Nef (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression, Th1

References Wasik et al. 2000

- HIV+ infants that progressed rapidly to AIDS had lower Th1 responses and decreased production of beta-chemokines and IL-2 relative to other HIV+ infants.
- No HIV+ infants had no demonstrable CTL at birth, but Th1 responses accompanied by CTL responses developed in children with slowly progressive disease, and not in rapid progressors.
- CTLp frequencies were determined by limiting dilution using autologous B cells infected with vaccina/HIV constructs.

**HXB2 Location** Nef

**Author Location Nef** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References De Maria et al. 1997

- CD3+ cells that also carry a natural killer cell receptor (NKR+) can exhibit down regulation of T cell function.
- Anti-NKR IgM MAb masked this inhibitory function and increased HIV-1 specific CTL activity in phytohemagglutininactivated PBMC cultured in the presence of IL-2 from 3/5 patients, and in one other case anti-NKR MAb brought HIV-1 specific CTL activity to detectable levels.

**HXB2 Location** Nef

**Author Location Nef** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Lubaki et al. 1999

- Three strategies were used to analyze CTL activity: area under the net HIV-specific lysis curve (ACU), linear regression (LR) of net specific lysis, and the standard method, lytic units (LU20)
- A correlation between low HIV plasma viral load and increased levels of HIV-specific Gag and Nef CTL activity was observed using ACU and LR, but not LU20.

**HXB2 Location** Nef

Author Location Nef (LAI)

Epitope Subtype B

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade LAI, B clade SF2 HIV component: Env, Gag, Nef, Protease

Species (MHC) human

References Gorse et al. 1999b

- The vaccine used was rec canarypox expressing HIV-1 env, gag, pol, nef and protease (vCP300) with or without administration of HIV-1 SF-2 rgp120.
- In vitro inducible CTL activity against HIV-1 Env, Gag, Pol, and Nef antigens was observed in 79% (15 of 19) of vaccine recipients.

• The combination of vCP300 and vP1291 together resulted in an overall increase in CTL induction and detection sensitivity.

**HXB2 Location** Nef

**Author Location** Nef

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords TCR usage

References Gamberg et al. 1999

- 13/13 subjects with advanced HIV infections showed CD8 T cell proliferation and differentiation of CTL in vitro, and six individuals showed HIV-specific responses to Gag, Pol, Env or Nef antigens.
- Data suggests that the functional and genetic integrity of the CD8 T cell repertoire (TCR betaV gene intrafamily genetic diversity) remains intact through advanced HIV infection, although HIV-specific CTL activity decreases.

**HXB2 Location** Nef

**Author Location** Nef

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Nef,

Rev, Tat

Species (MHC) human

Keywords HAART, ART

References Calarota et al. 1999

- 9/9 HIV-1 + subjects were given one of three DNA vaccinations for nef, rev or tat, and novel proliferative and CTL responses were generated.
- The nef DNA immunization induced the highest and most consistent CTLp activity, IFN-gamma production, and IL-6 and IgG responses.
- Highly active antiretroviral treatment (HAART) did not induce new HIV-specific CTL responses but reduced viral load, while DNA vaccination induced new immune responses but did not reduce viral load – thus this is a potentially complementary and promising combination.

**HXB2 Location** Nef

**Author Location** Nef (LAI)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Buseyne et al. 1998a

• This study showed a correlation between strong CTL memory and breadth of response in 7-12 month old infants, and remaining AIDS-free for the first year of life, higher absolute CD4 and CD8 cells, and lower viral load.

**HXB2 Location** Nef

**Author Location** Nef (LAI)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons **References** Buseyne *et al.* 1998b

 In infants with positive CTL responses, most responses showed cross-clade reactivity with somewhat diminished recognition of epitopes from different subtypes.

HXB2 Location Nef

**Author Location** Nef (LAI)

Epitope Subtype B Immunogen vaccine

*Vector/Type:* canarypox *HIV component:* Gag, gp120, gp41, Nef, Protease, RT

Species (MHC) human

References Evans et al. 1999

 A Canarypox vaccine expressing gp120, gp41, Gag, Protease, Nef and Pol CTL epitopes gave rise to CTL that could be detected in 61% of the volunteers – responses to Gag, Env, Nef and Pol were detected 3-6 months after the last vaccination.

**HXB2 Location** Nef **Author Location** Nef

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References da Silva & Hughes 1998

 CTL dense regions of Nef tend to lie in conserved domains with low non-synonymous substitution per site – authors consider that this may be due to a host adaptation to infection that focuses the CTL response to be directed against conserved functional domains da Silva & Hughes [1998]

**HXB2 Location** Nef

Author Location Nef (LAI)

Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Legrand et al. 1997

- Seventeen recently infected patients were tested for CTL response to HIV proteins Env, Gag, Pol, Rev, Nef, Vif and Tat.
- An early response (within a month following PI) was noted in 87% of the subjects to Gag, 75% to Env, and 50% to Nef.
- Early responses to Pol, Rev, Vif and Tat were rare.

**HXB2 Location** Nef

Author Location Nef (LAI)

Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Zerhouni *et al.* 1997

CTL responses to Env, Gag, Nef and RT were tested at various phases of disease progression – 10 asymptomatic patients generally had CTL responses to all proteins, 10 ARC patients responded well to all proteins except Nef, and AIDS patients had few responses to any proteins.

HXB2 Location Nef Author Location Nef Epitope Immunogen HIV-1 infection Species (MHC)

**Keywords** epitope processing **References** Kuiken *et al.* 1999

- A correlation between conserved regions of Nef and CTL epitope density was also noted in Kuiken et al. [1999]. The authors suggest that this may be due to biological reasons such as the one described above da Silva & Hughes [1998], or due to epitope processing, or may be an artifact of experimental strategy for epitope definition, such that conserved epitopes would tend to be identified because they are more likely to be cross-reactive with the test reagents.
- Both p17 and Nef show a correlation between epitope density and conserved regions in the protein; in contrast, p24 is a more conserved protein, and known epitopes are evenly distributed across p24.

**HXB2 Location** Nef

Author Location Nef (BRU)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Aladdin *et al.* 1999

 In vitro measurements of CTL-activity by Cr release assay in bulk culture showed no correlation between CTL-activity (gp120, Gag, Pol and Nef) and disease progression as measured by viral load, CD4 and time to death.

**HXB2 Location** Nef

**Author Location** Nef (SF2)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Jin et al. 1998a

 CTL precursor frequencies were determined in HIV-1 infected pregnant women, and higher CTLp frequencies to Pol and SF2 Nef, but not IIIB Nef, were found in non-transmitting mothers than in transmitting mothers – Nef CTL responses have been found in uninfected infants born to HIV+ women (Lazuriaga95);

**HXB2 Location** Nef

**Author Location** (subtype C)

Epitope Subtype C

Immunogen

Species (MHC) human

**Keywords** subtype comparisons, immunodominance **References** Novitsky *et al.* 2001

- This study is provides a survey of CTL responses and full length HIV-1 genome sequences from a C subtype infected Botswanan cohort.
- 37 of 45 subjects (82%) demonstrated Nef specific ELISPOT CTL responses of more than 100 SFC/106 PBMC.
- Two Nef-immunodominant regions were identified, one spanned amino acid positions 67 to 96 using HXB2 numbering system while the second corresponded to amino acid positions 122 to 141.

While there was some subtype B and C cross-reactivity, there was greater breadth and intensity of response if the CTL from HIV-1-infected individuals was probed with ELISPOT using peptides derived from the same subtype (a median of three Nef epitopes recognized within subtype C compared with one Nef epitope recognized from subtype B peptides, and ELISPOT results with a median of 763 SFC/106 PBMC among responses to HIV-1 C, versus a median of 318 SFC/106 PBMC among responses to HIV-1 B.

**HXB2 Location** Nef

Author Location Nef (subtype A, B, D)

**Epitope** 

Subtype A, B, D

Immunogen HIV-1 infection

Species (MHC) human

Keywords subtype comparisons

References Cao et al. 2000

- HIV-1 subtypes A and D dominate the Ugandan epidemic, and a vaccine trial using B clade antigen is underway – this study addresses relative levels of cross-reactive CTL responses in HIV infected Ugandans to A, D, and B clade recombinant vaccinia viruses expressing Gag, Env, Pol, RT or Nef from HIV-1 clades A, B, and D.
- Proteins corresponding to the subtype of the infecting strains tended to trigger higher levels of CTL response measured by percent specific lysis, but there was extensive inter-subtype cross-reactivity with B clade proteins and the co-circulating subtype.

**HXB2 Location** Nef

**Author Location** Nef

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA HIV component: Nef, Rev, Tat Adjuvant: CpG immunostimulatory sequence (ISS)

Species (MHC) human

Keywords review

References Calarota & Wahren 2001

 This review discusses the cellular immune response, and comments on the stimulatory role of CpG motifs and how HIV-1 DNA vaccines can boost the CTL and Th proliferative responses in asymptomatic HIV+ individuals.

**HXB2 Location** Nef

**Author Location** 

**Epitope** 

**Subtype** B **Immunogen** HIV-1 exposed seronegative

Species (MHC) human

**Keywords** HIV exposed persistently seronegative (HEPS), responses in children, mother-to-infant transmission

References De Maria et al. 1994; Kuhn et al. 2002

- 6/24 HIV uninfected infants (ages 15-50 months) born to HIV+ mothers had HIV-1 specific CTL responses to vacciniaexpressed Nef, Gag/Pol, Env.
- Reviewed in Kuhn et al. [2002].

**HXB2 Location** Nef

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** epitope processing, escape

References Yusim et al. 2002

- Epitopes that have been described in the literature and included in this database tend to cluster in conserved regions and be absent or rarely found in highly variable regions in Nef, Env, and p17.
- While the uneven distribution of epitopes may be in part due to a limited cross-recognition of specific responses because of differences between peptides used to probe the immune response and autologous strains, regions with a paucity of defined epitopes also had higher frequencies of amino acids that tend to not be found in C-terminal positions of epitopes, and had lower cleavage prediction scores for epitope processing. This suggests that in the regions of the virus where variation is best tolerated, traces of immune escape have left an imprint on the viral population. Epitopes also were concentrated in alpha-helix and turn regions in the proteins.
- In the more conserved p24, and Pol proteins RT and Protease, epitopes are more evenly distributed.

**HXB2 Location** Nef

Author Location Nef (HXB)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** epitope processing, vaccine-specific epitope characteristics

References Lu et al. 2000a

 Bacillus anthrax lethal toxin (LFn)-HIV fusion proteins are candidate HIV vaccines that are safe in mice, and LFn-V3 region fusion proteins induce CD8 T cells in BALBc mice. LFn causes exogenous protein to be taken up and processed in a class I pathway. Expressed proteins from Gag p24 and nef fragments cloned into the LFn expression plasmid stimulate gag-specific CD4 proliferation and CTL responses in HIVinfected donor PBMCs in vitro.

**HXB2 Location** Nef

**Author Location (BRU)** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Edwards *et al.* 2002

- 96% (26/27) chronically infected HIV-1 infected patients elicited gamma-IFN CD8+ T-cell responses against Gag.
- Nef and/or Pol CTL responses were detected in 86% of the subjects.
- The magnitude and breadth of Gag and p24 T-cell responses correlated with absolute CD4 counts, and inversely correlated with viral load.
- Pol and Int CTL responses correlated positively with absolute CD4+ T-cell count.

or viral load.

**HXB2 Location** Nef **Author Location Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, dendritic cells

References Larsson et al. 2002b

· Autologous mature dendritic cells with rec vaccinia expressing Gag, Pol, Nef and Env could amplify CD8+ T-cell Elispot responses 4-38 fold in five HIV+ patients on successful HAART treatment, relative to autologous monocytes. Some weak responses could only be detected using mature dendritic cells as APCs, and this approach could be useful for detection of low frequency memory cells.

**HXB2 Location** Nef **Author Location (SF2) Epitope** Subtype B

Immunogen HIV-1 and HCV co-infection

Species (MHC) human

**Keywords** rate of progression References Lauer et al. 2002

- HIV-1 and HCV immune responses were studied in 22 individuals who were co-infected with HIV-1 and hepatitis C virus (HCV). IFNgamma production was measured in an Elispot assay of CD8+ T-cells using targets expressing either Gag, RT, Env and Nef in a vaccinia construct, or one of seven HCV
- All 22 patients targeted at least one protein. 20/22 patients recognized RT, 17/22 patients recognized Gag, 13/22 subjects recognized Env and 11/22 patients recognized Nef. Robust CTL activity was independent of disease progression or viral load.
- Despite high HCV viral loads, very few HCV CD8+ T-cell Elispot responses were detected. In a control HCV infected person who did not have HIV-1, strong anti-HCV responses were mounted.
- HIV-specific CD4 proliferative responses were detected in 9/17 coinfected patients, but no HCV responses were detected.

**HXB2 Location** Nef **Author Location Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, responses in children

References Scott et al. 2001

- CTL responses before and after initiation of ART were studied in 13 HIV-1 vertically infected infants <6 months of age, and 4 that were >6 months of age.
- Before ART 2/13 infants <6 months of age showed IFNgamma Elispot CD8+ T-cell responses, one to Nef and one to Env and Nef, and these responses became undetectable after successful therapy-3 infants were coinfected with CMV and all 3 had CMV-specific CD8+ T-cell responses.

- Nef and Env responses did not correlate with either CD4 counts One older infant, at 23 months, had CTL responses against all for proteins tested, Gag, Pol, Nef and Env, and had the lowest plasma viremia of the study group. 3/4 infants older than 6 months of age responded to either Nef or Pol.
  - · Administration of ART over 48 weeks broadened the HIV-1specific CTL response in 2/4 of the older children that were incomplete responders.

**HXB2 Location** Nef

**Author Location (IIIB)** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, supervised treatment interruptions (STI)

References Ortiz et al. 2001

• Immune responses in eight chronically HIV-1 infected patients undergoing HAART therapy structured treatment interruptions (STI) were studied. STI boosted HIV-1 specific CTL responses and elevated CTL responses were maintained up to 22 weeks after the last treatment interruption, but viral load rebound to pretreatment levels and CD4 T-cell count decline was observed. CD8 responses in PBMC were measured by cytokine flow cytometry with gp160, Gag p55, RT-Pol and Nef expressed in vaccinia.

**HXB2 Location** Nef

**Author Location** 

**Epitope** 

Immunogen vaccine

Vector/Type: adenovirus HIV component: Gag-Pol, Nef, Vpr

Species (MHC) mouse

References Muthumani et al. 2002

- Vpr can cause cells to go into G2 arrest, and it surpresses immune cell activation and inflammatory cytokine production, so co-immunization of BALB/c mice with recombinant adenovirus expressing Vpr and HIV-1 antigens Nef or Gag/Pol was tested to see if Vpr reduced the immune response to the other HIV antigens.
- Vpr compromised CD8+ T-cell lytic responses and T-helper proliferative responses in mice co-immunized with Vpr and Nef or Gag/Pol.
- In vitro, Vpr reduced T-cell cytokine production of IL-12 and TNFalpha, indicative of Vpr-mediated immune suppression.

**HXB2 Location** Nef

**Author Location Nef** 

**Epitope** 

Subtype multiple

Immunogen

Species (MHC) human

Assay type Flow cytometric T-cell cytokine assay

Keywords subtype comparisons

References Currier et al. 2003

CD8-cellular immune responses from 21 HIV-1 infected patients from Kenya infected with subtype A, C, D, and unique recombinants were studied for cross-recognition of Gag, Env, and Nef vaccinia-expressed proteins representing subtypes A-H, including CRF01.

- Both subtype-specific and cross-reactive CTL were observed, with a skewing of responses towards the infecting subtype in the nine subjects with full length sequence available. The magnitude of the responses to Gag were the highest, less to Nef, and still less to Env.
- For Gag, 8/21 subjects responded to at least 7/8 different subtype proteins, 7 had a mixed response, recognizing some subtypes but not others, and 6 responded to only one or none of the different subtype proteins. For Env, 4/19 subjects responded to at least 7/8 subtypes, 8 had a mixed response, and 7 responded to one or none. For Nef, 5/19 subjects responded to at least 7/8 subtypes, 8 had a mixed response, and 6 responded to one or none.

**HXB2 Location** Nef

Author Location Nef (B.AU.AF064676)

Epitope Subtype B Immunogen

Species (MHC) human

References

HXB2 Location Nef
Author Location Nef
Epitope
Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot

- IFNγ, Intracellular cytokine staining **Keywords** assay standardization/improvement

References Draenert et al. 2003

- Six different HIV-1 Nef peptide sets ranging in length from 15-20 amino acids with overlap from 10-11 amino acids derived from either the B clade consensus sequences or the B clade sequence B.AU.AF064676 were used to study the impact of using different peptide design strategies to detect CD4 and CD8 T-cell responses. 20 individuals were tested using the six sets of peptides. 17/20 had CD8 T-cell responses to all peptides sets, and 15 of these 17 had CD4 T-cell responses.
- Although there was a trend in detecting more CD8 T cell responses using the shorter 15-mer peptides, longer 20-mers were best for detecting more CD4 T-cell responses, but neither result was statistically significant. Similar results were seen in the 15 to 20 amino acid range for both IFN gamma Elispot and ICS assays.
- Use of the consensus versus the natural strain identified slightly increased numbers of reactive peptides. Seven reactive peptides were observed with the B consensus peptides but not the B.AU.AF064676 peptides, but on the other hand four reactivites were observed using the B.AU.AF064676 peptides but not the consensus.
- Using an overlap of 10 or 11 amino acids did not make a difference.

HXB2 Location Nef

**Author Location** (C consensus)

**Epitope** Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords rate of progression

References Novitsky et al. 2003

 In this study, PBMC from 105 asymptomatic HIV-1 C clade infected patients from Botswana were screened for HIV-1 subtype C specific T-cell responses directed against Gag, Pol, Vif, Vpr, Tat, Rev, Vpu, Env and Nef. Nef-specific T-cell responses positively correlated with plasma viral load. In contrast, HIV-1 Gag and especially Gag p24 showed an inverse correlation with viral load.

**HXB2 Location** Nef

Author Location (C consensus)

**Epitope** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression

References Novitsky et al. 2003

 In this study, PBMC from 105 asymptomatic HIV-1 C clade infected patients from Botswana were screened for HIV-1 subtype C specific T-cell responses directed against Gag, Pol, Vif, Vpr, Tat, Rev, Vpu, Env and Nef. Nef-specific T-cell responses positively correlated with plasma viral load. In contrast, HIV-1 Gag and especially Gag p24 showed an inverse correlation with viral load.

**HXB2 Location** Nef

**Author Location Nef** 

**Epitope** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Country South Africa.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** rate of progression, immunodominance, acute/early infection, early-expressed proteins

References Masemola et al. 2004a

- Anti-HIV T-cell responses in subtype C HIV-1 infected individuals in the beginning of the infection target multiple protein regions, but the responses are dominated by Nef, making up almost one-third of the total responses. 97.5% of the Nef epitopes targeted were within a short stretch of 119 amino acids.
- Neither breadth nor magnitude of CD8+ T-cell responses were correlated with control of virus, however hierarchical preferential targeting of Gag was significantly associated with lower viral loads.

**HXB2 Location** Nef

Author Location Nef (B consensus)

Epitope

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** HAART, ART, immunodominance, acute/early infection, vaccine antigen design

References Lichterfeld et al. 2004b

- HIV-1 specific CD8 T-cell responses in individuals with acute and early HIV-1 infection are preferentially directed against epitopes in the central region of Nef, with 94% of the magnitude of the response in acute infection directed at Nef, and 46% during early infection. In chronic infection, CD8 T-cell immune responses are broadly diversified towards Gag, Env and Pol, and Nef accounts for only 17% of the response.
- The region of Nef that is targeted is the central most conserved region, but relative to other HIV proteins it is still quite variable. However, responses are cross-reactive enough to detect strong acute responses using consensus based peptides, and is an early expressed gene so may have advantages in the context of a vaccine.
- Nef immunodominance was retained in patients that were treated during acute infection, but no treatment and so continuous antigen exposure resulted in rapid diversification of the immune response.

## II-B-24 HIV-1 CTL/CD8 + epitopes

**HXB2 Location** HIV-1

**Author Location** 

Epitope

Subtype CRF01\_AE

Immunogen vaccine

Species (MHC) human (A11)

**Keywords** review, vaccine-specific epitope characteristics, escape

References Ariyoshi et al. 2002

 This review summarizes a meeting held to discuss options for determining CTL responses to vaccines. Problems are noted: costs for some assays are prohibitive for a Phase III study, Elispot shows interlaboratory variation but could be extended to many samples. HLA-A11 is very common in Thailand – over 30% carry the HLA-A11 allele. Predominant strains may be evolving to evade recognition of A11 restricted epitopes. Few full length CRF01 sequences are available. Epitopes may differ in vaccinees and infected individuals.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human (A11, B8, B40, Cw8)

Assay type Cytokine production, CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining

**Keywords** HAART, ART, acute/early infection, early treatment

References Alter et al. 2003

• Longitudinal study (24 mo) monitoring T-cell immune responses in 4 patient groups: Group 1 (n=6) consists of subjects who underwent HAART preseroconversion, group 2 (n=11) were HAART treated during early postseroconversion, group 3

(n=5) contained patients who started HAART during late postseroconversion, and group 4 (n= 6) commenced with HAART during chronic HIV-1 infection.

- The experimental strategey was to test for reactivity levels with sets of peptides that each contain epitopes with known HLArestricting elements, making the peptide selection based on the optimal epitope list in this database. The HLA alleles found in the patients were balanced so that the frequency in the groups were comparable. Peptides spanning parts of Gag, Env, Nef, and RT were used for Elispot, and Gag peptides were used for ICS.
- All group 1 patients, and 5/11 group 2 patients, maintained the breadth and the magnitude of the immune response throughout the study; those in group 2 that maintained response started therapy earlier. The hierarchy of intensity of responses to different peptides was preserved. Individuals in groups 3 and 4 all showed a decline, and after treatment lost responses. Groups 1 and 2 showed HAART-induced suppression of viremia but maintained resposnes. Groups 3 and 4 both showed viral suppression in association with a decreased immune response in breadth and magnitude after HAART. The authors suggest that preservation of HIV CD4+ responses can be maintained even if HAART is first given beyond the acute phase of infection, and a delay may allow a full CD8 response to develop while still allowing CD4 function to be preserved.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Species (MHC) human (B27, B8)

**Keywords** binding affinity, review, subtype comparisons, epitope processing, escape

References McMichael & Hanke 2002

- CTL response-eliciting vaccines are reviewed. The natural epitope interactions with the HLA class I presenting molecules and T-cell receptors are described, and the impact of breadth of CTL responses and diversity considered in a vaccine context.
- Interesting specific examples are given concerning anchor chain residues. For B27, the B pocket fits Arg (R) but not Lys (K), so even this conservative change is not tolerated. In B8 either R or K can fit in the B pocket, but the substitution will cause conformational shifts in other parts of the epitope.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Vector/Type: Listeria monocytogenes HIV

component: Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords review

References Lieberman 2002

Attenuated Listeria monocytogenes vectors elicit strong persistent CTL responses in vaccinations of BALB/c mice and can protect mice from a vaccinia-gag challenge.

HXB2 Location HIV-1

**Author Location** gp120 (V3) and p24 (IIIB, MN, BH10)

**Epitope** 

**Subtype** A, B **Immunogen** vaccine

Vector/Type: virus-like particle (VLP) Strain: A clade UG5.94UG018, B clade IIIB

HIV component: Gag, gp120

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type Chromium-release assay Keywords subtype comparisons References Buonaguro *et al.* 2002

- Different HIV strains were used for different regions: gp120 A clade UG5.94UG018, and B clade IIIB
- BALB/c mice were given intraperitoneal immunization with virus-like particle (VLPs) expressing recombinant subtype A gp120 and Pr55gag in the absence of adjuvants.
- High dose-independent humoral responses against both gp120 and p24 peptides were detected. Antibodies able to elicit 50% neutralization against A clade IIIB and the autologous clade a virus were obtained.
- Recombinant rgp120 (clade B, MN) induced T-cell proliferative responses *in vitro* from vaccinated animals.
- CTL activity was observed against splenocytes expressing Env (clade A) and Gag (clade B, BH10) from a vaccinia construct.

**HXB2** Location HIV-1

**Author Location** 

Epitope

Subtype B

Immunogen vaccine

Vector/Type: DNA, polyepitope Strain: A clade, B clade HIV component: Env, Gag, Pol Adjuvant: IL-12, IL-2, liposome

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type CD8 T-cell Elispot - IFNγ, Intracellular cytokine staining, Delayed-type hypersensitivity

(DTH), Chromium-release assay

**Keywords** vaccine-induced epitopes **References** Shinoda *et al.* 2004

 Mice immunized with a polyepitope DNA vaccine encoding 20 antigenic epitopes of several HIV-1 clades (hDNA vaccine) showed strong Ab responses, activation of IFN-gamma secretion cells targeting gp120 and synthetic antigenic peptides, and several peptide specific CTL responses. When challenged with recombinant HIV-vaccinia viruses, mice immunized with the hDNA vaccine showed lower viral titers in the ovary.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Schito et al. 2001

 Longitudinal analysis (72 weeks) of 15 patients with acute or recent HIV-1 infection implies that HAART treatment alone can not completely conserve CD8+ cell homeostasis and preserve the original T-cell receptor repertoire.

HXB2 Location HIV-1 Author Location Epitope Immunogen HIV-1 infection

Species (MHC) human

References Mackewicz et al. 2000

 Non-cytotoxic anti-HIV responses of CD8+ T cells cultured with CD4 infected HIV cells are mediated be blocking expression of viral RNA, and do not influence viral replication steps through integration of provirus.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Species (MHC)

Keywords dynamics

References Altes et al. 2002

- This study employs a mathematical model to study the consequences of increasing the T-helper response through a vaccine, which would have counter-balancing effects in a new infection: a more intense response provides more help but also more target cells. The model indicates that if the infecting virus had a low replication rate, then CTLp and CD4 helper cells could control an infection. Only a vaccine that could increase CTL responsiveness could reduce viral set point with observed replication rates.
- A CD4+ T-cell response without maintained CTL response was deleterious in this model.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords assay standardization/improvement

References Currier et al. 2002b

- Elispot standardization was sought using a reference peptide pool of 23, 8-11 mer epitopes from Influenza, cytomegalovirus (CMV), and Epstein Bar Virus (EBV) presented by 11 common HLA class I molecules.
- 15/17 (88%) HIV- and 14/20 (70%) HIV+ individuals reacted with this test set and *in vitro* simulation of the PBMC from these individuals were capable of killing cells expressing the target antigen.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human, macaque

Keywords dynamics, HAART, ART

References Wodarz 2002

 Mathematical modeling is used to support the idea that T-helper cell dysfunction results in a compromised ability to maintain an anti-HIV CTL memory response. Models suggest strategies to restore CTL memory through therapy and improve long-term immunological control of the virus.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Species (MHC) human

Keywords review

References Zinkernagel 2002

 HIV immunity and vaccine strategies are compared with to other pathogens. We do not have a successful vaccine agains TB leprosy, HIV, HCV and most parasites, and the author suggests this is associated with the need for a strong T-cell response to these diseases. Vaccine strategies that achieve a physiological low does infection that is well controlled but persists may be required to alter the immunopathological consequences of infection with HIV.

**HXB2** Location HIV-1

**Author Location** 

Epitope

Immunogen vaccine Species (MHC) human

**Keywords** review, subtype comparisons, epitope processing

References Gaschen et al. 2002

- The concept of using an artificial consensus sequence for vaccine design is discussed, comparing the concepts of a model ancestor sequence or a consensus sequence, with illustrations of the potential advantages of the strategy based on C-clade comparisons.
- See also a comment Nickle *et al.* [2003], and reply Gao *et al.* [2003]

**HXB2** Location HIV-1

**Author Location** 

Epitope

Immunogen HIV-1 infection Species (MHC) human, macaque

**Keywords** review, class I down-regulation by Nef, escape **References** Johnson & Desrosiers 2002

- Reviews evidence for CTL escape in HIV epitopes in natural human infections, and in SIV infections of macaque where viral clones with a known time of infection and multiple animals with the same HLA molecules can be tracked.
- Vigorous CTL responses are made despite class I downregulation by the Nef protein, but it may delay cytolysis of infected cells. Too great a loss of MHC proteins may enhance NK cell killing so the fitness advantage of this function of Nef may be in balance.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Species (MHC) human

**Keywords** review, epitope processing, supertype, com-

putational epitope prediction, HIV exposed persistently seronegative (HEPS), supervised treatment interruptions (STI), immunodomi-

nance

References Newman et al. 2002

- This extensive review covers many aspects of T-cell immunity and natural HIV infections, and considers how this knowledge might be applied to a polyepitope vaccine approach. Strategies concerning ways to avoid the creation of junctional epitopes and use of linkers to enhance processing of such constructs are discussed.
- The C-terminal flanking residue (C1) was found to be associated with immunodominance of epitopes, such that R or K (positive charge) > N or Q (amide) > C, G, A, T, S (small) > F, W, Y (aromatic) > I, L, M, V (aliphatic) > D (negative). As this position is outside and proximal to the epitope, processing and cleavage is the likely reason for this observation.
- Changing the C1 residue from F to K for an HLA-A2 presented epitope from HBV resulted in a change from the epitope being non-immunogenic to strongly immunogenic.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Species (MHC) human

**Keywords** review, HIV exposed persistently seronegative (HEPS)

References Johnston & Flores 2001

 Reviews the current state of HIV vaccine approaches, and discusses the role of CTL induced immunity in protection or partial protection in animal studies, likening it to the CTL found in HEPS studies.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords binding affinity, review, escape

References Klenerman et al. 2002

- The importance of breadth, or spread, of CTL responses is discussed, as narrowly focused responses can be more readily escaped.
- Some HLA types and specific epitope recognition may be associated with a better disease outcome. Reasons for this are considered, including NK cell activity, epitope affinity, epitope conservation, and class I specific induction of more effective T-cell receptors.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** review, HIV exposed persistently seronegative (HEPS), responses in children, mother-to-

infant transmission

References Kuhn et al. 2002

 Intrauterine exposure of infants to HIV from their mothers results in HIV-1 specific T-helper cell proliferative responses in 1/3 of exposed uninfected babies, and HIV-1 specific CTL in some. Such responses are evident, but it is unknown whether they are associated with lack of infection, but there is some evidence that HIV-1 T-cell responses may reduce transmission and CD8 responses detected in earlier studies.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords HIV exposed persistently seronegative (HEPS), mother-to-infant transmission

References Kuhn et al. 2002; Levy et al. 1998

- A non-HLA-specific, non-chemokine-mediated CD8+ T-cell non-cytotoxic anti-HIV response, measured by suppression of acute viral infection of CD4 cells, was detectable in approximately 16/31 (52%) of uninfected children born of infected mothers, was more commonly detected in those <1 year old, and could reflect a protective response.
- Reviewed in Kuhn et al. [2002].

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Species (MHC) human

Keywords dynamics

References Altes et al. 2001

• Mathematical modeling suggests if the effector CTL vaccine response exceeds the level of response seen in chronic infection, that a memory CTL population is established that can respond very quickly to protect from infection.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Species (MHC) human

Keywords review

**References** Copeland 2002

• This review summarizes cytokines and chemokines produced by CD8+ T-cells that can interfere with HIV's infection and replication.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Species (MHC)

Kevwords review

References Edgeworth et al. 2002

• This review summarizes HIV vaccine strategies, adjuvants, current clinical trials and animal models.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Species (MHC)

Keywords review

References Graham 2002

in breastfeeding mothers. Summary tables are provided of CD4 • This review summarizes HIV vaccine approaches and clinical

**HXB2 Location** HIV-1

**Author Location** Env (HXB2)

**Epitope** 

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade HXB2 HIV component: gp140ΔCFI, gp160 deletions

Species (MHC) guinea pig, mouse

References Chakrabarti et al. 2002

- Intramuscular injection of plasmid DNA was used to vaccinate BALB/c or Huntley guinea pigs with a series of codonoptimized modified HIV-1 HXB2 envelopes - modifications included elimination of glycosylation sites, deletions, and exchange of the V3 loop to change from a X4 or R5 phenotype.
- The mutant envelope gp140deltaCFI gave the most promising result, enhancing antibody responses while retaining the ability to stimulate a strong CTL response.
- gp140deltaCFI has deletions in the cleavage site, fusogenic domain and spacing of the heptad repeats, and was designed to mimic a fusion intermediate.

HXB2 Location HIV-1

Author Location Env (gp160) (384-467)

**Epitope** 

Immunogen vaccine

Vector/Type: hepatitis B surface antigen lipoprotein particles (HsBAg) Strain: B clade LAI HIV component: V3

Species (MHC) macaque, rabbit

References Michel et al. 1993

• Immunization with recombinant HIV1 V3/HBsAg hybrid particles into rabbits or macaques elicited and maintained for several months anti-V3 or HIV-1 Env proliferative, CTL and Ab responses.

HXB2 Location HIV-1

Author Location Gag (HXB2)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Garba et al. 2002

- CD8+ Tcells from 25% of HIV positive individuals produce TGF-beta1 in response to stimulation with HIV proteins, and this can significantly reduce CD8+ T-cell IFN-gamma induction to HIV and vaccinia proteins.
- Different peptides can preferentially induce TGF-beta1 or IFNgamma from CD8+ T-clymphocytes from the same individual, and TGF-beta1 non-specifically suppresses HIV-specific immune responses.

**HXB2 Location** HIV-1

**Author Location** Pol (HXB2)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Garba et al. 2002

- CD8+ Tcells from 25% of HIV positive individuals produce TGF-beta1 in response to stimulation with HIV proteins, and this can significantly reduce CD8+ T-cell IFN-gamma induction to HIV and vaccinia proteins.
- Different peptides can preferentially induce TGF-beta1 or IFN-gamma from CD8+ T-clymphocytes from the same individual, and TGF-beta1 non-specifically suppresses HIV-specific immune responses.

**HXB2 Location** HIV-1 **Author Location** Env (MN)

Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Garba et al. 2002

- CD8+ Tcells from 25% of HIV positive individuals produce TGF-beta1 in response to stimulation with HIV proteins, and this can significantly reduce CD8+ T-cell IFN-gamma induction to HIV and vaccinia proteins.
- Different peptides can preferentially induce TGF-beta1 or IFN-gamma from CD8+ T-clymphocytes from the same individual, and TGF-beta1 non-specifically suppresses HIV-specific immune responses.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining

**Keywords** assay stand

standardization/improvement,

acute/early infection

References Altfeld et al. 2003

- The frequency of HIV-1 specific T-cell responses was characterized in an Elispot IFN-gamma assay, using 507 overlapping peptides based on the B clade consensus sequence spanning all HIV-1 clade B proteins against PBMC from 57 HIV-1 infected patients at various disease and treatment stages. 63% of the peptides were recognized (range of 1-42 per subject, median=14). More variable peptides were targeted less frequently.
- Autologous virus sequences from six patients in acute infection spanning of HIV-1 p24, Tat and Vpr were used to scan for missed responses due to viral variation when using the consensus for peptides. 12/42 (29%) responses to these peptides were dectected only with autologous peptides, and often these autologous reponses were immunodominant. Responses were also generally higher using autologous peptides.
- A longitudinal analysis (5 yrs) of the T-cell responses in 5 patients showed that the autologous sequence detected stronger T-cell recongition than the HIV-1 clade B consensus sequence.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection Species (MHC) chimpanzee

Keywords review

References Balla-Jhagihoorsingh et al. 2003

- This paper reviews HIV-1-specific cell-mediated immune responses in chimpanzees and discusses mechanisms that might control HIV-1 pathogenesis in chimpanzees. During the first decade of the HIV epidemic, more than 200 chimpanzees were experimentally infected with HIV. Among these only one case of declining CD4+ cells has been reported, all others have remained asymptomatic with no loss of immune function, some after 20 years of infection. In contrast to infected humans which have a skewed Th2 response, chimpanzees maintain balanced Th responses and are likely to support a fully mature CD8+ T-cell response.
- Specific HIV epitopes recognized by chimpanzees have been mapped and CTL detected, but overall the responses are at much lower levels than in humans, as viral loads are so low. Gag epitope responses are estimated to be 0.0095 to 0.0025% of the CD8+ T cell population in chimpanzee, and 1-2% in humans.
- The authors argue that the chimpanzee immune response may
  be effective at controlling virus because it focuses on conserved epitopes, and further speculate that long contact with
  lentiviruses may have put strong selection pressures on the
  chimpanzee MHC class I, narrowing the population's ability to
  respond to only the most conserved, and so useful, epitopes.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** HAART, ART, supervised treatment interruptions (STI)

**References** Fagard *et al.* 2003

This study monitored the effects of repeated treatment interruptions (STI), in 2-week intervals, in 133 HIV-1 infected, HAART-treated patients. STIs were rarely able to control viremia without continued HAART, and increases in CD8+T-cell response frequencies did not correlate with the level of control of viral replication. CD8+T cell responses were measured by gamma IFN Elispot using between 2-32 different optimal HIV epitopes, selected to be appropriate for the patient's HLA type.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen

Species (MHC) human

References

HXB2 Location HIV-1

**Author Location HIV-1** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** responses in children **References** Feeney *et al.* 2003

- The magnitude and breadth of CD8+ T-cell responses in 18 pediatric (6-17 years) perinatally HIV-1 infected patients was determined using 1) overlapping peptides spanning all HIV-1 proteins and 2) peptides from all predefined appropriately class I HLA-restricted HIV-1 epitopes.
- Perinatally infected children's CD8+ T-cell responses were comparable in magnitude and breadth to adult responses. Many reactive peptides did not overlap with a previously characterized optimal epitope.
- On average 20% of all known pre-defined optimal epitopes presented by appropriate HLAs were recognized in these children. In two patients, autologous sequences spanning unrecognized potential epitopes usually corresponded to the reactive form of the epitope, so epitope variation alone did not account for unrecognized epitopes.
- · Children with detectable viremia showed a broader and greater CTL responses than HAART responsive children with undetectable viremia.

**HXB2 Location** HIV-1

Author Location HIV-1

**Epitope** 

Immunogen

Species (MHC)

References

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Species (MHC)

Kevwords review

References Hanke 2003

• Review of HIV vaccine development discussing diversity, the merits and difficulties of stimulating different arms of the immune response, and different strategies, including DNA vaccines, viral vectors, CTL epitope based, and protein- or peptidebased vaccines.

**HXB2 Location** HIV-1

**Author Location** HIV-1 (HXB2)

**Epitope** 

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC) human

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

Keywords HIV exposed persistently seronegative (HEPS)

References Hladik et al. 2003

- Longitudinal study analyzed IFN-γ CD8+ T cell responses in highly exposed, seronegative homosexual men. Overlapping peptides spanning the Gag, Env, Nef and Pol subtype B HXB2 sequence were used to stimulate PBMC from 26 individuals, whose frequency of HIV-1 specific IFN- $\gamma$ T cell responses were
- CD8+ T cells from 3/15 individuals (EES15, ES29, and ES63) recognized > 3 peptide pools.

**HXB2** Location HIV-1 **Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type Chromium-release assay

Keywords dynamics

References Kousignian et al. 2003

• The diversity of HIV protein (Gag, Pol, Env, Nef, Rev, Tat, Vif) recognition by CTLs was studied longitudinally in a cohort of 152 HIV-infected untreated individuals, and was analyzed by Markov modelling. CTL responses from 152 HIV-1 infected patients in four stages of disease progression were collected for a period of 5 years. Results show that memory CTL responses against HIV-1 proteins are acquired during early HIV-1 infection and subsequently lost. As viral load increased there was an accelerating loss of multiple protein recognition.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: gp120 depleted whole killed virus Adjuvant: Incomplete Freund's Ad-

juvant (IFA)

Species (MHC) human

References Lederman & Douek 2003; Robbins et al. 2003

• Lederman and Douek is an editorial comment referring to the study presented by Robbins et al., in which the authors discuss why an HIV-1 gp120-depleted inactivated HIV vaccine elicits HIV-1 specific T helper responses in 5/5 HIV+ people, but not CD8+ CTL responses. In chronically infected people it appears that stimulating Th responses in and of itself is not enough to restore strong CTL responses.

HXB2 Location HIV-1

**Author Location** 

Epitope

Immunogen vaccine

Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72), CpG immunostimulatory se-

quence (ISS), HSP70

Species (MHC) human

Keywords review, Th1, Th2, genital and mucosal immu-

References Lehner 2003

• This review discusses the importance of mucosal and innate immunity for future vaccination strategies in HIV infection in humans. Different mucosal adjuvants are compared, and the advantages of a Th1 polarized response.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Onyemelukwe & Musa 2002

Longitudinal study (1991-1997) of the clinicial presentation of 80 HIV-1 or HIV-II seropositive people in Zaria, Nigeria, who contracted HIV-1 primarily via heterosexual transmission. Main complicating diseases were tuberculosis and bacterial infections including Salmonella, Streptococcus peneumoniae

and Stahhylococcus. HIV-1 progression was associated with a decline of not only CD4+ T cells, but CD8+ T cells as well patients had CD4+ counts < 200 cells/ul, and CD8 counts were 190 cells/ul versus 440 cells/ul for controls.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**References** Onyemelukwe & Musa 2002

• Longitudinal study (1991-1997) of the clinicial presentation of 80 HIV-1 or HIV-II seropositive people in Zaria, Nigeria, who contracted HIV-1 primarily via heterosexual transmission. Main complicating diseases were tuberculosis and bacterial infections including Salmonella, Streptococcus peneumoniae and Stahhylococcus. HIV-1 progression was associated with a decline of not only CD4+ T cells, but CD8+ T cells as well patients had CD4+ counts < 200 cells/ul, and CD8 counts were 190 cells/ul versus 440 cells/ul for controls.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot

- IFNγ

Keywords HAART, ART

References Price et al. 2003

- CD4+ and CD8+ T cell responses were analyzed in this longitudinal study (19 mo) of 53 patients with chronic HIV-1 infection receiving continous ART therapy. Three subgroups were compared: one with suppressed viremia and increasing CD4+ T cell counts, one with dectable viral load and declining CD4, and one with detectable viral load with a positive CD4+ T cell slope.
- IFN-γ ELISPOT analysis was performed with peptides spanning RT, Env, Gag (p24), Gag(p17), Nef, Tat and Rev. The IFN-γ analysis showed the greatest CD4+ as well as CD8+ Tcell responses in the group with stable CD4+ T cell responses despite detectable virus over a median time course of 9 months.

**HXB2** Location HIV-1

**Author Location** 

**Epitope Immunogen** 

Species (MHC) human

Assay type Chromium-release assay

**Keywords** rate of progression

References Sindhu et al. 2003a; Sindhu et al. 2003b

• In a cross-sectional study of 31 HIV+ people, a correlation was observed between CTL-mediated bystander HLA-unrestricted lysis of primary CD4+ T-cells.  $\gamma\delta$  CTL are abnormally expanded in HIV+ people, and the V $\delta$ 1 subset can deplete bystander CD4+ T-cells and expedite progression. In a set of 13 patients, an inverse correlation was observed between CD8+ Tcell activation markers and viral load, thought to be an indicator of CTL-associated immunopathogenesis in HIV progression.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC)

Keywords review

**References** Vella & Daniels 2003

• This article reviews the CD8+ T-cell antiviral factor (CAF). CAF contributes to MHC restricted, CD8+ T-cell mediated non-cytolytic suppression of HIV in infected individuals.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Subtype A, B, C

Immunogen vaccine

Vector/Type: DNA, polyepitope HIV component: gp120, gp41, Nef, p17 Gag, p24 Gag, Pol Adjuvant: concavalin A-immobilized

polystyrene nanospheres

Species (MHC) mouse

Assay type proliferation, CD8 T-cell Elispot - IFN $\gamma$ 

Keywords vaccine-induced epitopes

References Bazhan et al. 2004

• A synthetic T cell polyepitope immunogen containing 80 overlapping Env, Gag, Pol and Nef epitopes was used to immunize mice. It induced both humoral and cellular responses which increased upon reimmunization.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

**Immunogen** 

Species (MHC)

**Keywords** review, class I down-regulation by Nef, earlyexpressed proteins, immune evasion

References Collins 2004

• There are a number of factors that combined make HIV-infected cells resistant to CTLs. HLA-associations with disease progression are reviewed. Nef down-regulation of HLA class I A and B molecules is one important mechanism of HIV immune evasion. Rev allows late viral proteins to be expressed, enabling CTL specific for epitopes in these proteins to recognize infected cells. It is suggested that blocking the activity of Nef and Rev would reduce production of viral variants and enhance the ability of CTLs to combat HIV.

HXB2 Location HIV-1

**Author Location (SIV)** 

**Epitope** 

Immunogen SIV infection

Species (MHC) macaque

Keywords escape, reversion, viral fitness

References Friedrich et al. 2004

• SIV CTL escape variants revert to wild-type epitopes after transmission to new hosts with disparate MHC class I alleles. Thus mutations in CTL epitopes may have moderate to severe negative effects on viral replicative fitness although some escape variants are shown to accumulate substitutions in flanking regions of the epitope that help compensate for fitness loss.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen

Species (MHC) human

Keywords dynamics, HAART, ART

References Ganusov 2003

 The rate of virus decline after initiation of HAART is shown by a mathematical model, to depend on whether the virus is controlled by the CTL response via lytic or non-lytic mechanisms.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** review, class I down-regulation by Nef, escape, dendritic cells, TCR usage, memory

cells, immune dysfunction

References Gulzar & Copeland 2004

 HIV has developed numerous strategies to evade CD8+ T-cell response that are reviewed in this paper, including escape mutations in CD8+ T-cell recognition, down-regulation of MHC-I surface expression, alternating cytokine production, disruption of proper CD8+ T-cell signaling resulting in anergy, and disruption of the function of CD4+ T-cells and APCs required for CD8+ T-cell maturation.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

**Immunogen** in vitro stimulation or selection

Species (MHC) human

**Assay type** CD8 T-cell Elispot - IFNγ, Chromium-release assay, Flow cytometric T-cell cytokine assay

**Keywords** characterizing CD8+ T cells

References Kitchen et al. 2004

 This paper characterizes a population of cells that are CD3+, CD8+, and CD4+. These cells are mature and highly activated. The CD4 molecule expressed by these CD8+ T-cells plays an important role in expression of IFN-gamma and Fas ligand and cytotoxic responses. HIV infection of CD8+CD4+ T-cells results in Nef independent down-regulation of CD4 and dysregulation of IFN-gamma and Fas ligand, and provides an additional resevoire for the virus.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen

Species (MHC) human

Keywords review, characterizing CD8+ T cells

References Petrovas et al. 2004

This review discusses the attributes of HIV-specific CTLs that
contribute to their inability to control HIV infection, with
an emphasis on the susceptibility of HIV-specific CTL to
CD95/Fas induced apoptosis upon binding target cells. Furthermore, Nef may inhibit apoptosis by blocking CD95/Fas
signaling on infected cells.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Country** United Kingdom.

Assay type proliferation, CD8 T-cell Elispot - IFNγ, T-

cell Elispot, Flow cytometric T-cell cytokine

assav

Keywords HAART, ART, immune dysfunction

References Pires et al. 2004

Daily administration of rec human growth hormone (rhGH) induced an increase in the numbers of naive CD4 T-cells and effector CD8 T-cells. Also, a rise in HIV-1 antigen-specific CD4 and CD8 T-cell responses was observed. The function of specific effector CD8 T-cells was perserved despite an eventual decrease of specific CD4 T-cell responses.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Species (MHC) human

Country United States.

Assay type CD8 T-cell Elispot - IFNγ

**Keywords** review, supervised treatment interruptions (STI), vaccine-specific epitope character-

istics, variant cross-recognition or cross-

neutralization

References Robinson 2003

This paper is a commentary on Altfeld et al, Nature 420:434
2002. The patient AC-06 was superinfected with a second
strain of HIV-1 after STI despite 12 of 25 recognized CD8+ Tcell epitopes maintaining strong cross-reactive immunity measured by gamma IFN EliSpot against the second strain. While
vaccine trials in macaques have given optimistic results, this
patient's superinfection in spite of a strong cross-reactive CD8+
T-cell immune response suggests that vaccine strategies may
have to be re-examined.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human

Country Spain.

Assay type Flow cytometric T-cell cytokine assay

**Keywords** rate of progression

References Rodés et al. 2004

A complex set viral or host factors has been found to be associated with the absence of disease progression among long-term non-progressors (LTNP). 19 LTNP were followed for six years; 12 were non-progressors over this period, 7 showed a slow progressive CD4 depletion. Their virus replicative capacity was shown to be reduced and T-cell activation was low. Pooled peptide CD8+ T-cell gamma IFN responses did not differ between non-progressors, slow progressors, or a group of HIV progressors.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** Subtype B

Immunogen HIV-1 infection, vaccine

Vector/Type: canarypox prime with recombinant protein boost Strain: B clade SF2 HIV component: Env, Gag, Protease

Species (MHC) human

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ

**Keywords** assay standardization/improvement

References Russell et al. 2003

• IFNγ Elispot assay is shown to be a good initial screening method for measurment of CD8+ T-cell responses in both vaccination and natural HIV-1 infection. Responses were detected using peptides at low concentrations (1-2 µg/mL) and an increase in detection of HIV-1 specific CD8+T-cells by using 15-mers rather than 20-mer peptides for cell activation was observed. More responses were detected using smaller pools (10 or 2 peptides) than larger pools (25 or 50 peptides), so smaller pools may be needed to detect low frequency responses. Responses to natural infection were more than a log higher than to the vaccine.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, SIV infection

Species (MHC) human, macaque

Keywords review, escape, reversion, viral fitness

References Smith 2004

• This paper reviews several studies which track HIV and SIV CTL escape mutations after transmission into a new host, and reversion rates and fitness costs of CTL escape. Some escape mutants have a cost to viral fitness. The author suggests that CTL based HIV-1 vaccine should therefore not only increase cellular responses against viral epitopes but also favor epitopes where escape mutations result in significant decrease in viral fitness.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** Subtype B Immunogen vaccine

> Vector/Type: DNA Strain: B clade HIV component: gp140, gp160 Adjuvant: re-

ovirus alpha 1 protein

Species (MHC) mouse

Donor MHC H-2d

Assay type Cytokine production, Chromium-release assav

Keywords adjuvant comparison

References Wang et al. 2003

• M cells are found in the follical-associated epithelium in mucosal inductive tissues, and reovirus are able to attach to these cells via the alpha 1 protein. Respiratory mucosal sites were targeted with a reovirus protein alpha 1 protein delivered with a DNA vaccine administerd i.n. in BALB/c mice. The naked gp160 DNA vaccine did not elicit CD8+ T cell reponses, but when delived with alpha 1 protein, CTL responses were observed in the lungs, spleens and lymph nodes. gp160 was shown to be most immunogenic compared to a cytoplasmic gp140 and secreted gp140. The vaccinated animals had reduced vaccinia virus when challeneged with a vaccinia-env recombinant.

HXB2 Location HIV-1

**Author Location** Gag

**Epitope** 

Immunogen vaccine

Vector/Type: DNA with CMV promotor, fowlpoxvirus Strain: SIV HIV component: Env, Gag, Pol, Rev, Tat, Vpu Adjuvant: IFNγ, CpG immunostimulatory sequence (ISS)

Species (MHC) macaque

Assay type proliferation, T-cell Elispot, Intracellular cytokine staining

Keywords adjuvant comparison, vaccine antigen design References Dale et al. 2004

Macaques immunized with DNA and fowlpox vaccines showed high levels of CD4 and CD8 T-cell immune responses to Gag. Single DNA priming vaccination or coexpressed IFN-gamma with the fowlpox virus boost were shown to be less immunogenic and less protective than sequental DNA and fowlpox virus vaccination. Partial protective immunity was observed following a high dose, virulent SHIV challenge, for the DNA fowlpox prime boost, as well as the DNA vaccination alone, even though standard assays failed to detect a strong immune respsonse with DNA alone.

**HXB2 Location** HIV-1

Author Location (IIIB, Thai B', Chinese CB)

**Epitope** 

Subtype B, C

Immunogen HIV-1 infection

Species (MHC) human

Country China.

Assay type Intracellular cytokine staining, Chromiumrelease assay

**Keywords** subtype comparisons, characterizing CD8+ T cells

References François-Bongarcon et al. 2004

• The ability of circulating T-cells from 7 North American and 4 Chinese HIV+ donors to produce IFN-gamma and/or lyse autologous primary cells infected with HIVIIIB, B' (Thai B) or C/B recombinant form was tested. The results showed crossclade CD8 T-cell responses to the Cinese viruses among North American donors and to HIVIIIB in Chinese donors, suggesting that many of the T-cell responses to clade B virus epitopes are conserved across clades. Lysis of cells by N. American donor CD8+ T cells infected with IIIB or a Thai B' strain were comparable, while lysis infected with the Chinese BC recombinant was somewhat reduced, although the reduction was not statistically significant.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type Cytokine production, Chromium-release assay, Flow cytometric T-cell cytokine assay

Keywords TCR usage, memory cells, characterizing

CD8+ T cells, immune dysfunction

References Gamberg et al. 2004b

• A relationship was found between the proportion of HIVspecific CTLs expressing CD28 and CD4+ T-cell counts, viral load and disease progression. This association cannot be linked to disease related degeneration of CD8+CD28- T-cells in terms of their TCRbetaV family repertoire diversity or ability to produce cytokines. This suggests that effective immune responses contain CD8+CD28+ T-cell populations that shift to CD8+CD28- in ineffective responses.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen

Species (MHC)

Keywords review, epitope processing, vaccine-specific epitope characteristics, rate of progression, immunodominance, escape, acute/early infection, early-expressed proteins, TCR usage, reversion, viral fitness

References Goulder & Watkins 2004

• CTLs have a central role in the control of HIV infection. Emergence of escape variants to CTLs is one of the major obstacles to vaccine development. Factors that should be considered for the development of an HIV vaccine are CTLs that are specific for epitopes recognized during the acute phase of infection, CTLs that are able to efficiently control viral replication, and epitopes from regions of the viral genome that are highly conserved or where variation results in loss of viral fitness.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

Country Kenya.

Assay type Chromium-release assay

Keywords HIV exposed persistently seronegative (HEPS), characterizing CD8+ T cells

References Kaul et al. 2004

• HIV-1 specific CTL responses found in HIV exposed persistently seronegative Kenyan female sex workers were shown to be associated with age and recent HIV-1 exposure, but not with protection against HIV-1 infection. The authors note that CTL may be the result of a non-productive HIV infection, but not mediate protection; alternatively, the low incidence, possibly due to behavioral interventions, may not give adequate sampling to detect the response.

**HXB2** Location HIV-1 **Author Location Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Country France.

Keywords HAART, ART, characterizing CD8+ T cells,

immune dysfunction

References Kryworuchko et al. 2004

• A subset of HIV-1 infected untreated patients had CD8+ T-cells that were unable to respond to IL-2 by activating STAT5a and b proteins. This was correlated with an impaired activation of the upstream kinase Jak-3. 6 months of HAART was shown to restore Jak/STAT signalling in those patients and their CD8+ T-cell response to IL-2. This suggests another mechanism for immune dysfunction in HIV infected patients.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Subtype A, B, C, D, F, G, U

Immunogen computer prediction

Species (MHC) human

Keywords vaccine antigen design

References Maksyutov et al. 2004

• Every HIV protein was shown to have some regions that were highly similar to the regions of human proteins. Most of those regions contained T-cell or/and B-cell epitopes. The epitopes shared by HIV and its host may have immunopathogenic potential through stimulating autoimmunity and should possibly be excluded from HIV vaccines. All HIV proteins from the sequence of BH10 were compared to human proteins, as well as many HIV-1 V3 variants.

**HXB2** Location HIV-1

**Author Location (ELI)** 

**Epitope** 

Immunogen in vitro stimulation or selection

Species (MHC) human

Assay type Cytokine production, proliferation,

Chromium-release assay, Flow cytometric

T-cell cytokine assay Keywords class I down-regulation by Nef, rate of pro-

gression, dendritic cells, immune dysfunction

References Quaranta et al. 2004

• Exogenous Nef protein activates immature DCs and inhibits the capacity of DCs to prime CD8+ T-cell responses by downregulating their proliferation and function capacities. Nef induces CD8+ T-cell apoptosis by up-regulating TNF-alpha and FasL production by DCs, while DCs are protected from apoptosis themselves. These mechanisms, as well as by down regulation of the HLA class I proteins, can contribute to HIV-triggered immune dysfunction.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Switzerland.

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ

**Keywords** rate of progression **References** Oxenius *et al.* 2004a

In untreated, HIV-1 chronically infected patients, CD4+ T-cell responses and, to a lesser extent, CD8+ T-cell responses, were found to inversely correlate with disease progression rate. Polymorphisms in CCR genes, HLA genotype and GB virus C coinfection were not found to be related to slower disease progression.

HXB2 Location HIV-1 Author Location (B clade)

> Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type CD8 T-cell Elispot - IFN $\gamma$ , Flow cytometric

T-cell cytokine assay

Keywords genital and mucosal immunity, characterizing

CD8+ T cells

References Sheth et al. 2004

• HIV viral load in semen is found to be 10-fold lower than in blood. No correlation was found between viral load in either semen or blood and systemic HIV-specific CD8 T-cell responses, in 20 samples.

HXB2 Location HIV-1

**Author Location** 

Epitope Subtype B

Immunogen vaccine

Vector/Type: DNA with CMV promotor, virus-like particle (VLP), modified vaccinia Ankara (MVA) Strain: B clade HIV component: Env, Gag, Pol, Protease, RT

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining

Keywords vaccine antigen design

References Smith et al. 2004

 Macaques were immunized with codon-optimized Gag DNA and non-codon-optimized Gag-Pol-Env DNA vaccines, expressed as VLPs as aggregates, followed by an MVA boost. There was no significant difference in anti-Gag T-cell responses and anti-Env Ab responses between the different vaccines. A second MVA boost did not increase T-cell responses but it increased anti-Env Ab titers by 40- to 90-fold.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope Immunogen** 

Species (MHC) human

Keywords review, epitope processing, rate of progres-

sion, escape, early-expressed proteins, vac-

cine antigen design

References Yang 2004

- This review considers CTL biology in HIV infection in the context of vaccine design principles. Since HIV-1 infection damages immunity through depletion of CD4+ T-cells, which in turn results in diminished capacity of the immune system to produce new and functional CTL responses, maximizing the breadth of CTL responses might not be enough for an HIV-1 vaccine. CTLs recognizing early proteins might be more prone to epitope escape mutation, while those recognizing more conserved structural proteins might be more likely to persist, so focusing on more conserved proteins those might be a good strategy to produce an attenuating vaccine.
- Original antigenic sin is discussed, the initial responses to an
  antigen that persist even after escape occurs, blunting the later
  immune response. If the goal is to prevent disease, focusing
  on conserved late expressed proteins might be the best target,
  where the fitness cost is greatest for escape; if the goal is to
  prevent infection, focusing the vaccine on the more variable
  early expressed proteins that elicit the first responses, Tat and
  Nef, might be best.

**HXB2** Location HIV-1

Author Location p24 (HIV-2 ROD, HIV-1 IIIB)

**Epitope** 

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human

Country Gambia.

Assay type Cytokine production, proliferation, CD8 Tcell Elispot - IFNγ, Chromium-release assay

**Keywords** rate of progression **References** Jaye *et al.* 2004

 A comparison of T cell responses in HIV-1 and HIV-2 infected asymptomatic patients with CD4+ cell counts 20% showed no significant difference between groups. Viral loads were roughly 20 times greater in HIV-1 positive patients than HIV-2 positive

patients.

• 10/20 (50%) of HIV-1 infected patients demonstrated proliferative responses with SI greater than 1.4 to gp120, and 11/20 to p24. 8/29 (29%) of HIV-2 infected patients recognized gp105, and 8/29 (29%) p26. Cytokine responses in both groups did not differ.

 9/21 (43%) of HIV-1 + and 15/30 (50%) of HIV-2 + patients had cyotoxic T cell responses to Gag, and 3/21 (14%) HIV-1 + and 8/30 (27%) HIV-2 + responded to Pol.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Country Spain.

Assay type proliferation, Intracellular cytokine staining

**Keywords** HAART, ART

References López et al. 2004

 A clinical trial compared chronically HIV-1 infected patients who had replaced HAART with didanosine (ddI) and hyroxyurea (HU) were followed for 12 months to an untreated HIV+ group and a group that continued on HAART.

- Approximately 20% of the patients treated with ddI-HU had detectable CD4+ T-cell proliferative responses to Gag and Env in contrast to drug-naive and HAART treated HIV-infected patients, who had few or no responses.
- HIV-specific CD8+ T-cell responses were higher in ddI-HU treated patients than HAART treated patients, even in individuals that maintained undetectable viral loads.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen

Species (MHC)

Keywords review, adjuvant comparison

References Mitchison & Sattentau 2005

 Review summarizes mechanisms of immunoregulation relevant for new vaccine development, with a brief summary of adjuvant triggering innate immunity through Toll-like receptors (TLRs), Nod molecules, and other activators. DNA encoded adjuvants that have been tested in DNA vaccines are summarized. The balance between Th1 (CTL activating) and Th2 (B cell activating) responses is discussed, and it is noted that BALB/c mice are predominately Th2 responders, C57BL Th1.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human Keywords escape

References Piontkivska & Hughes 2006

• Greatest amino acid diversity is found in sites in the HIV genome that are spanned by antibody epitopes. Sites spanned by CTL epitopes, but not by antibody epitopes, showed reduced amino acid diversity, even in comparison to non-epitope sites. However, mutations within CTL epitopes were more likely to be convergent than mutations within antibody epitopes. These patterns were consistent both in Gag and in Env.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFN $\gamma$ , CD8 T-cell Recy-

cleSpot - IFN $\gamma$ 

Keywords assay standardization/improvement, optimal

epitope

References Bihl et al. 2005

• This study describes a novel approach to achieve maximal information from an extensive set of antigens (HIV, EBV, CMV, HCV, and HBV) to determine the magnitude of T-cell responses while requiring minimal cell numbers. Large sets of peptides based on optimally defined epitopes from each pathogen are used. It is shown that, when compared to ex vivo cell preparations, antigen-unspecific in vitro T-cell expansion maintains the breadth of detectable T-cell responses. Also, harvesting cells from negative ELISpot wells for re-use (RecycleSpot) maximizes the use of available cells.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

 $\textbf{Keywords} \ \ \text{genital and mucosal immunity, characterizing}$ 

CD8+ T cells

References Ibarrondo et al. 2005

- The breadth and magnitude of HIV-1-specific CTL responses in blood and sigmoid colon mucosa were assessed in 16 patients. The magnitude of pool-specific CTL responses in blood and mucosa was correlated within each individual and across all individuals. CTL targeting was also found to correlate between these 2 compartments, with Nef being the most highly targeted region, followed by Gag. No correlation between the magnitude and breadth of CTL responses and viremia and blood CD4 levels was found in any of the compartments. Concordant peptide pool responses were found in the blood and mucosa 85%; pools that differed were near the threshold of detection.
- This study suggests that HIV-1-specific CTL responses in the blood mirror those in the mucosa during chronic infection.

HXB2 Location HIV-1

**Author Location** (Z321)

**Epitope** 

Subtype A, B, C, G

Immunogen vaccine

Vector/Type: gp120 depleted whole killed virus Strain: AG recombinant HZ321 HIV component: gp120 depleted virus Adjuvant: CpG immunostimulatory sequence (ISS)

Species (MHC) mouse

Assay type Cytokine production, proliferation, CD8 T-

cell Elispot - IFN $\gamma$ , Flow cytometric T-cell

cytokine assay

**Keywords** subtype comparisons, genital and mucosal immunity, adjuvant comparison, vaccine antigen

design, characterizing CD8+ T cells

References Jiang et al. 2005

- Mice were given intranasal immunization with inactivated gp120-depleted HIV-1 antigen plus a CpG ODN adjuvant and examined for local immune responses in the genital tract. Mice immunized with HIV Ag plus CpG produced significantly higher levels of IFN-gamma and beta-chemokines than mice immunized with Ag alone, and their lymphocites showed significant HIV-specific proliferation. CD8 T-cells were increased in the genital tracts of mice immunized with HIV Ag plus CpG.
- The vaccine antigen Z321 is clade G in Gag. Cross-clade protection against an intravaginal IVAG challenge was observed for clades A, C, and G, but not clade B.

HXB2 Location HIV-1

**Author Location** 

Epitope

Immunogen computer prediction

Species (MHC) human

Assay type Other

**Keywords** assay standardization/improvement, computational epitope prediction

References Larsen et al. 2005

 A computational epitope identification method integrating predictions of MHC class I binding affinity, TAP transport efficiency and C-terminal proteasomal cleavage was compared to two already existing computational epitope identification tools. It was shown that the new ANN method performed better, reducing the number of nonamers needed to be tested in order to identify 85% of the epitopes from 9-10% to only 7%.

HXB2 Location HIV-1

Author Location (A, B, and C consensus)

**Epitope** 

Subtype A, B, C

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD8 T-cell Elispot - IFN $\gamma$ 

**Keywords** subtype comparisons

References Yu et al. 2005

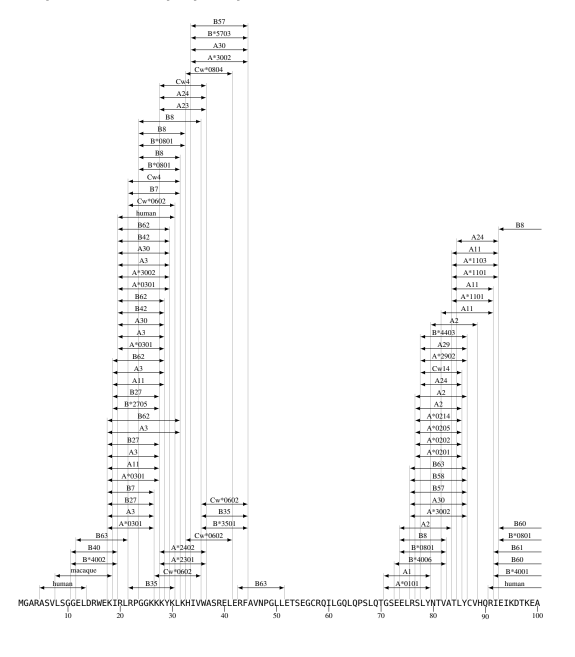
- HIV-specific T-cell responses to peptide from A, B, and C clade spanning the entire HIV proteome were assessed in clade B infected individuals. Many cross-reactive responses were observed against clade A, B, and C consensus sequences, preferentially recognized in conserved regions with low intra-clade diversity and high inter-clade homology.
- At the individual peptide level, within clade responses to B clade peptides were more frequent. 194 responses were detected with only one peptide, of these 105 recognized B clade, 55 C clade, and 34 A clade. 125 responses recognized peptides from two clades, and 110 of these were with B plus either A or C. 166 responses were cross-reactive with all three clades.

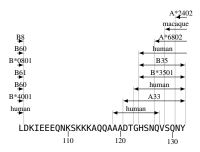
#### II-C

# Maps of CTL/CD8 + Epitope Locations Plotted by Protein

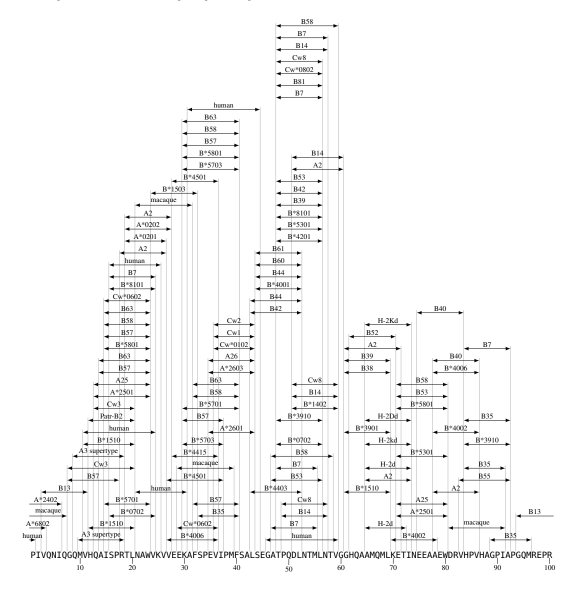
Linear CTL epitopes mapped to within a region of 14 amino acids or less are shown.

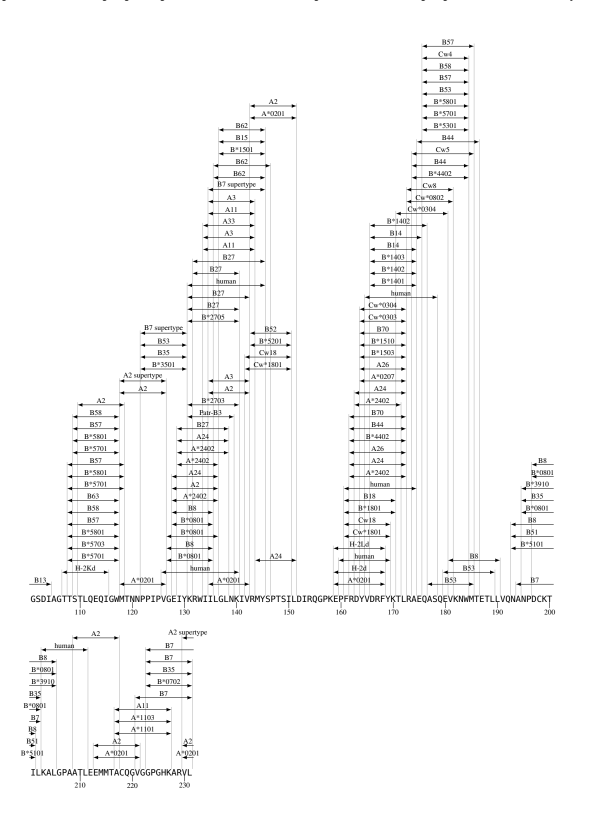
#### II-C-1 p17 CTL/CD8 + Epitope Map



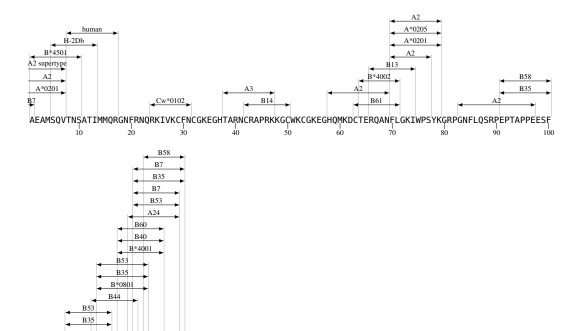


#### II-C-2 p24 CTL/CD8 + Epitope Map





#### II-C-3 p2p7p1p6 CTL/CD8 + Epitope Map



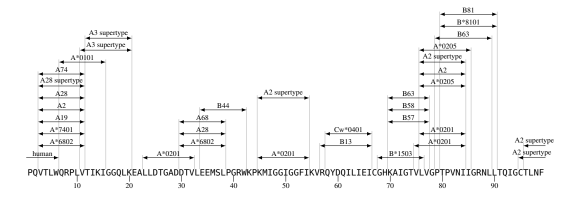
#### Gag/Pol TF CTL/CD8 + Epitope Map

RSGVETTTPPQKQEPIDKELYPLTSLRSLFGNDPSSQ 120

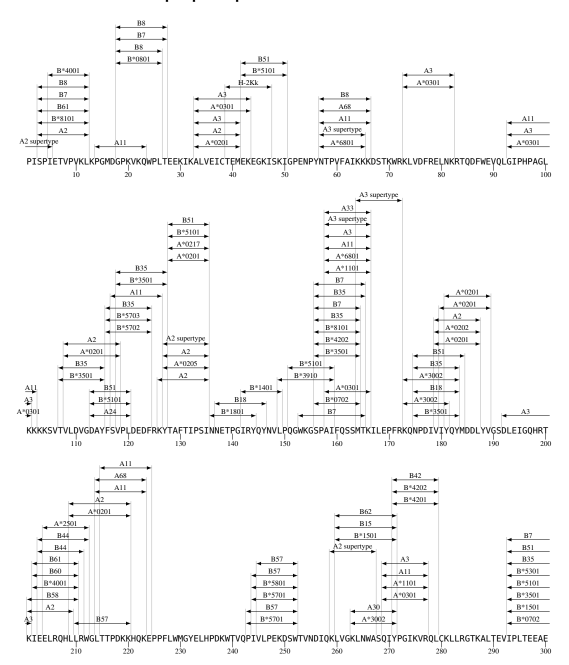
110

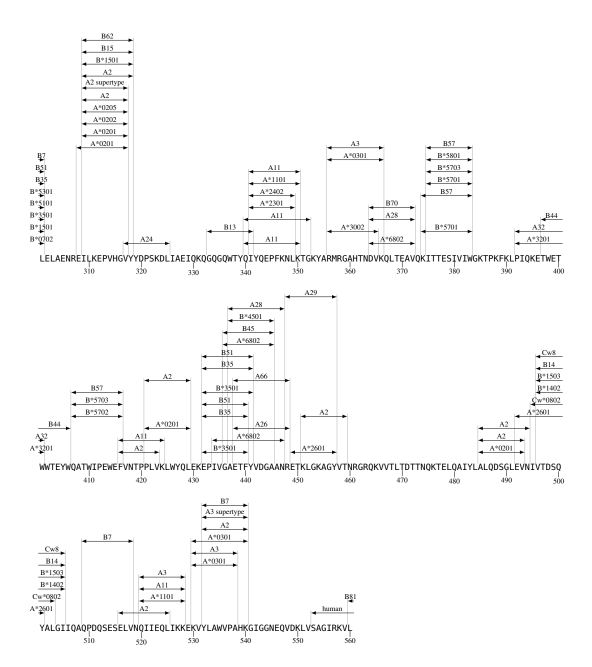


#### **Protease CTL/CD8 + Epitope Map**

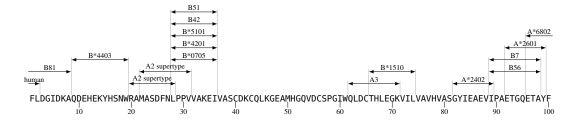


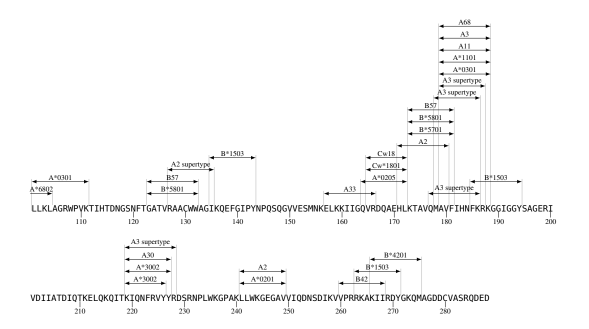
#### II-C-6 RT CTL/CD8 + Epitope Map



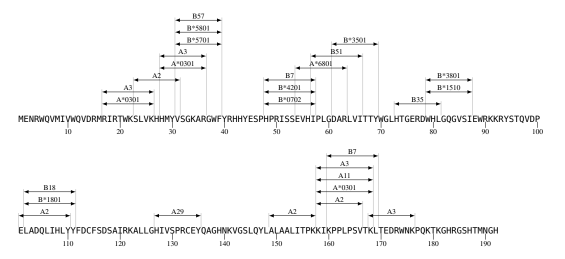


#### II-C-7 Integrase CTL/CD8 + Epitope Map

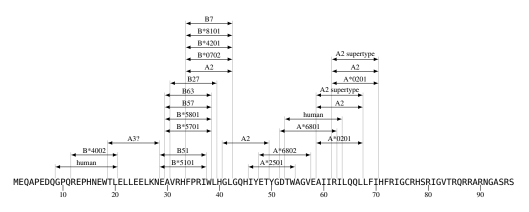




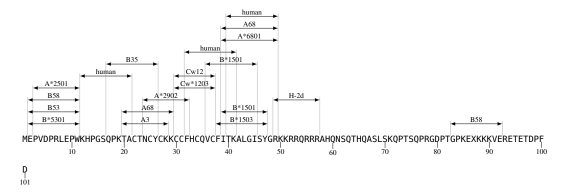
#### II-C-8 Vif CTL/CD8 + Epitope Map



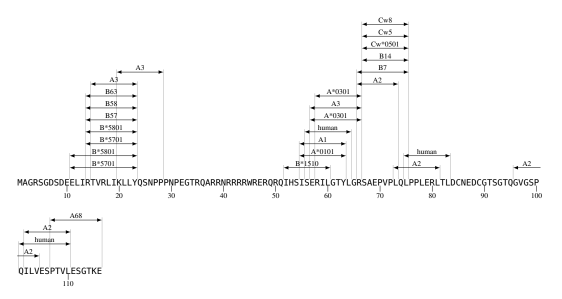
#### II-C-9 Vpr CTL/CD8 + Epitope Map



#### II-C-10 Tat CTL/CD8 + Epitope Map



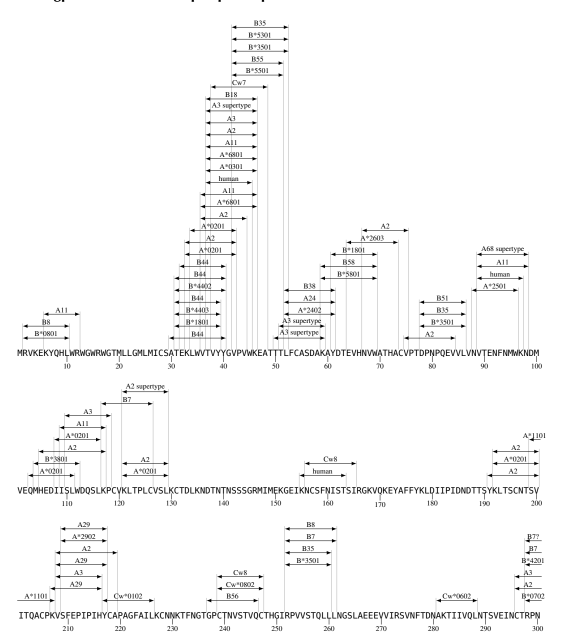
#### II-C-11 Rev CTL/CD8 + Epitope Map

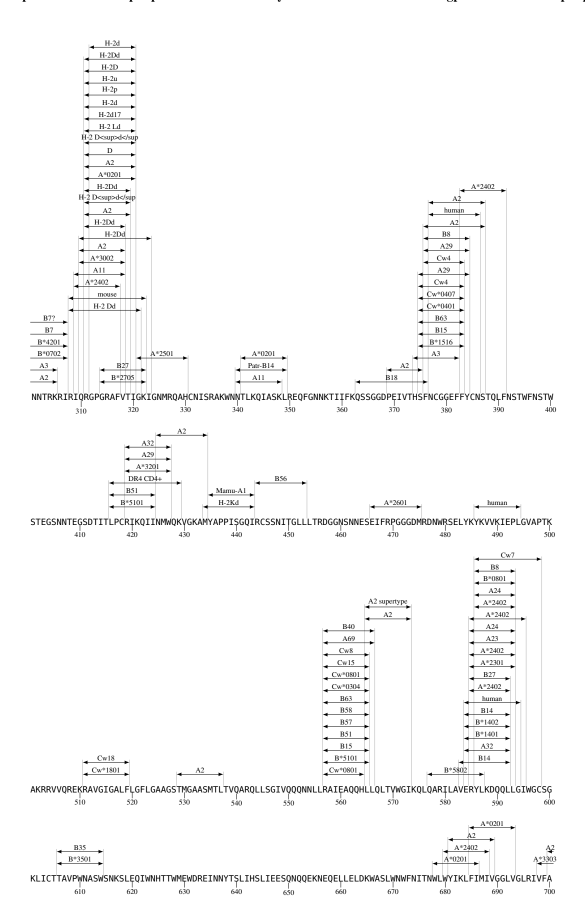


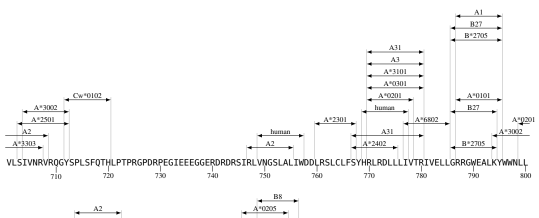
#### II-C-12 Vpu CTL/CD8 + Epitope Map

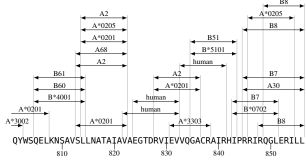


#### II-C-13 gp160 CTL/CD8 + Epitope Map

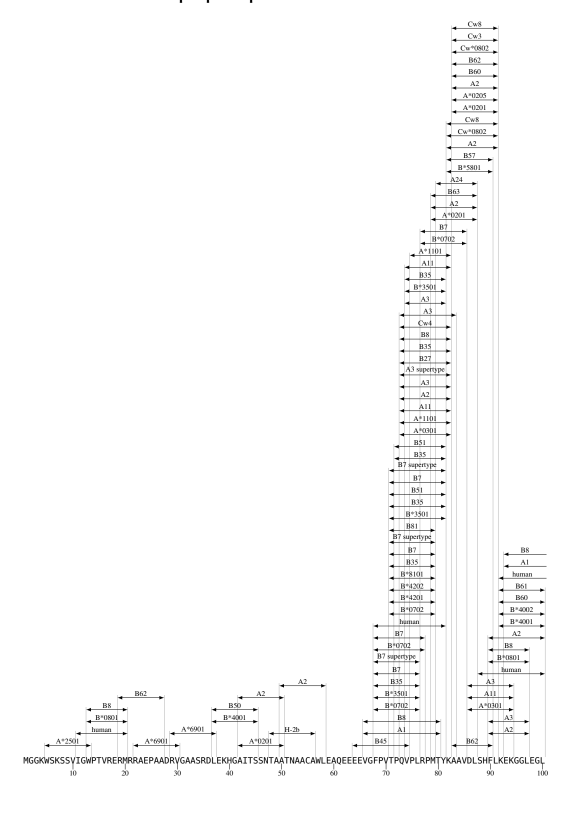


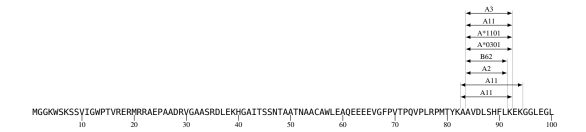


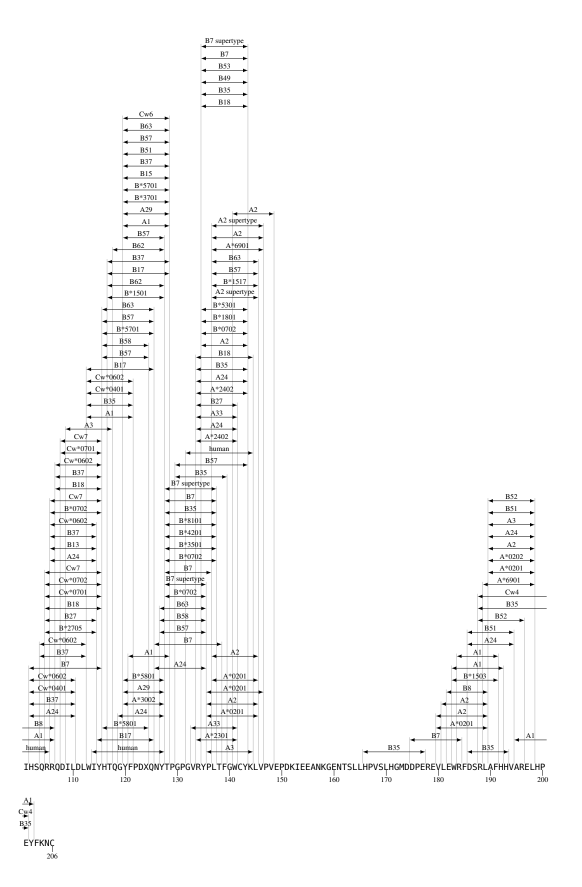




#### II-C-14 Nef CTL/CD8 + Epitope Map







# Part III

#### III-A

### **Summary**

This part includes tables, maps, and associated references of HIV-specific helper T-cell (Th) epitopes arranged sequentially according to the location of the proteins in the HIV-1 genome. We attempted to make this part as comprehensive as possible, requiring that the epitope be contained within a defined region of a maximum of 30 amino acids, but not that the optimal boundaries be defined. Studies that were based on the analysis of whole proteins are described at the end of each protein section. The same epitope can have multiple entries, as each entry represents a single publication in this part of the database. HLA specificity is usually not determined for Th epitopes. For more recent updates, epitope sequence alignments, and useful search capabilities, please see our web site: http://www. hiv.lanl.gov/content/immunology. Helper T-cell responses to proteins with no defined epitopes are listed at the end of each protein section.

Recent studies utilize multiple functions attributed to T cells to define responses, and the simple distinctions of cytotoxic T cells and helper T cells have become blurred as more is learned about the range of responses triggered in CD4 and CD8 positive T cells responding to antigenic stimulus. When adding the most recent studies to the database, we have tried to place T cell responses in a reasonable manner into our traditional helper T cell and CTL parts, and to specify the assay used to measure the response in each study.

#### **III-A-1** Epitope tables

Each T-helper epitope has a multi-part basic entry:

HXB2 location: The viral strain HXB2 (GenBank Accession Number K03455) is used as a reference strain throughout this publication. The position of the defined epitope location relative to the sequence of the HXB2 protein is indicated. The numbering in this table corresponds to the protein maps. Because of HIV-1 variation the epitope may not actually be present in HXB2; rather, the position in HXB2 indicates the position aligned to the epitope. HXB2 was selected as the reference strain because so many studies use HXB2, and because crystal structures for HXB2-related proteins are often available. The precise positions of an epitope on the HXB2 reference strain can be readily obtained using the interactive position locator at our web site: http://www.hiv.lanl.gov/

content/hiv-db/LOCATE/locate.html.

Author location: The amino acid positions of the epitope boundaries and the reference sequence are listed as given in the primary publication. Frequently, these positions as published are imprecise, and do not truly correspond to the numbering of the sequence, but they provide a reasonable guide to the peptide's approximate location in the protein. Also, in many cases the reference sequence identification was not provided, and in such cases it is not possible to use these numbers to specify precise locations.

Epitope: The amino acid sequence of the epitope of interest as defined in the reference, based on the reference strain used in the study defining the epitope. On occasions when only the position numbers and not the actual peptide sequence were specified in the original publication, we tried to fill in the peptide sequence based on the position numbers and reference strain. If the sequences were numbered inaccurately by the primary authors, or if we made a mistake in this process, we may have misrepresented the binding site's amino acid sequence. Because of this uncertainty, epitopes that were not explicitly written in the primary publication, that we determined by looking up the reference strain and the numbered location, are followed by a question mark in the table.

**Epitope name:** If the epitope has a name attributed by the publication, it is recorded here, e.g. "SL9".

**Subtype:** The subtype under study, generally not specified for B subtype.

**Immunogen:** The antigenic stimulus of the Th response. Often this is an HIV-1 infection. If a vaccine was used as the original antigenic stimulation, not a natural infection, this is noted separately, and additional information about the vaccine antigen is provided as available.

**Species (MHC):** The species responding and MHC or HLA specificity of the epitope.

**Donor MHC:** The HLA genotype of the individual that responded to the epitope.

**Country:** The country where the samples were obtained—generally not specified if the study was conducted in the United States.

**Assay type:** Assay used to characterize the response.

**Keywords:** Keywords are a searchable field for the web interface that is included in the T-cell sections of the printed version to help identify entries of particular interest.

**Reference:** The primary reference (sometimes two or more directly related studies are included). Details for some of the earlier references are in Part V.

Following the entry for a given Th epitope brief comments explain the context in which the epitope was studied and what was learned about the epitope in a given study.

#### III-A-2 HIV protein epitope maps

All HIV Th epitopes mapped to within a region of 18 amino acids or less are indicated on the HIV protein epitope maps. The location and HLA restriction elements of Th epitopes are indicated on protein sequences of HXB2. These maps are meant to provide the relative location of defined epitopes on a given protein, but the HXB2 sequence may not actually carry the epitope of interest, as it may vary relative to the sequence for which the epitope was defined. Epitopes with identical boundaries and HLA fields are included in the maps only once. If one laboratory determines HLA presenting molecules at the serotype level (example: A2) and another at the genotype level (example: A\*0201) both will be included in the map. MHC specificities are indicative of the host species; when no MHC presenting molecule is defined, the host species is noted.

#### **III-A-3** Alignments

To conserve space, no epitope alignments are provided in this book, but they can be generated using the Th epitope search tool at http://www.hiv.lanl.gov/content/immunology. The master alignment files from which the epitope alignments were created are available at our web site at http://www.hiv.lanl.gov/content/hiv-db/ALIGN\_CURRENT/ALIGN-INDEX.html.

#### III-B

### **HIV Helper/CD4+ T-Cell Epitope Tables**

All HIV Helper/CD4+ T-Cell epitopes are arranged by protein position. The table entries are sorted in a nested way—first by protein, then by HXB2 start location within the protein, and finally by HLA presenting molecule. Epitopes for which the HXB2 location is unknown appear at the end of the listing of the protein in which they are located.

## III-B-1 Gag p17 Helper/CD4+ T-cell epitopes

**HXB2 Location** p17 (1-18)

Author Location p17 (1-18 B consensus)

**Epitope MGARASVLSGGELDRWEK** 

Subtype B

**Immunogen** HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This epitope was recognized by 14% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location p17 (7–17)

Author Location Gag (7–17)

Epitope VLSGGELDRWE

Epitope name Gag 1.2

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT, Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining

Keywords subtype comparisons, memory cells

References Amara et al. 2005

- Clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better-conserved for CD8 than CD4 T cells.
- 3/5 CD8 epitopes and 2/8 CD4 epitopes were conserved across multiple HIV-1 clades. All 5 CD8, and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation. VLSGGELDRWE was not reported for human infections.
- The response elicited to the B clade epitope VLSGGEL-DRWE does not cross-react with the CRF02\_AG form VLtGGELDsWE. The forms VLSGG[e/k]LD[r/ak]WE are prevalent among M group clades.

**HXB2 Location** p17 (9–26)

Author Location p17 (9-26 B consensus)

Epitope SGGELDRWEKIRLRPGGK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 22% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were

seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location p17 (13–23) Author Location Gag (13–23) Epitope LDRWEKIRLRP Epitope name Gag 1.3

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT, Tat, Vpu

Species (MHC) macaque

**Assay type** T-cell Elispot, Intracellular cytokine staining **Keywords** subtype comparisons, memory cells

References Amara et al. 2005

- Clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better-conserved for CD8 than CD4 T cells.
- 3/5 CD8 epitopes and 2/8 CD4 epitopes were conserved across multiple HIV-1 clades. All 5 CD8, and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation.
- · LDRWEKIRLRP was not reported for human infections.
- The response elicited to the B clade epitope LDR-WEKIRLRP does not cross-react with the CRF02\_AG form LDsWEKIRLRP. Other clades most commonly carry an A in this position, and C clade consensus carries K (LD[r/ak]WEKIRLRP).

**HXB2 Location** p17 (17–31)

Author Location Gag (17–31 HXB-2)

Epitope EKIRLRPGGKKKYKL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** DRB1\*1303, DRB1\*1502, DRB3\*0101,

DRB5\*0102, DQB1\*0301, DQB1\*0601 **Country** United States.

Assay type CD4 T-cell Elispot - IFNγ

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 1/22 patients responded to this peptide.

HXB2 Location p17 (17-34)

**Author Location** p17 (17–34 B consensus)

Epitope EKIRLRPGGKKKYKLKHI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

692

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 14% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p17 (17–34)

**Author Location** p17 (17–34)

Epitope EKIRLRPGGKKKYKLHKI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production

References Geels et al. 2006

- The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.
- Autologous sequences corresponding to known and predicted Th epitopes were analyzed. EKIRLRPGGKKKYKLHKI had fixation of 1 mutation (EKIRLRPGGKK[k/r]YKLHKI) in 1 of the patients.

**HXB2 Location** p17 (18–42)

**Author Location** p17 (18–42 PV22)

Epitope KIRLRPGGKKKYKLKHIVWASRELE

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*13)

Donor MHC A29, A30, B8, B35, DRB1\*03, DRB1\*13

Keywords HAART, ART, Th1, Th2

References Lotti et al. 2002

- 10/49 chronically HIV-1 infected patients had low p55-Gagspecific CD4+ T cell responses prior to therapy, and these responses remained unchanged 3 and 6 months after initiation of HAART. There was no difference in level of response in those with or without a detectable p55 response.
- For one individual, patient F45 CDC stage A2, CD4+ p55 responding clones were generated. Her response was consistently strong and heterogeneous in terms of HLA restriction and  $V\beta$  usage, and some clones had a Th1 cytokine secretion profile

(high IFNgamma production) while some had a Th2 profile (high IL-4 and IL-5 production). 5/10 CD4+ clones could also induce cytotoxicity.

• 4/10 clones from patient F45 had their epitopes mapped using overlapping peptides. Clone 6 recognized this peptide sequence restricted by DRB1\*13. This clone had a high SI (27.1 to p55, 90.6 to peptide) secreted IFNgamma, indicative of a Th1 response, as well as TNFalpha. Clone 6 was highly cytotoxic, through a perforin-mediated pathway.

HXB2 Location p17 (21-35) **Author Location** p17 (21–35 SF2) Epitope LRPGGKKKYKLKHIV Immunogen HIV-1 infection Species (MHC) human (DR13.02) Keywords escape

References Harcourt et al. 1998

- 43 asymptomatic HIV+ individuals were screened for proliferative responses to HIV – 12 showed a response, and dominant epitopes were mapped for two individuals, one in p24 and one in p17.
- Patient 024's naturally occurring variant LRPG-GKKKYQLKHIV also elicited a strong proliferative • The most HIV-1 specific CD4+ T-cell responses were observed
- Naturally occurring variants of this epitope were found within the individual who made this response - several did not stimulate the CD4+ T-cell line that recognized the index peptide, suggestive of immune escape.

HXB2 Location p17 (22-29) Author Location p17 (22–29 LAI) Epitope RPGGKKKY? Subtype B Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

- Stimulates T-cell proliferation in HIV-infected donors.
- Schrier lists this peptide as p24(22-29), but it appears to be in p17.

HXB2 Location p17 (29-43) **Author Location** Gag (29–43 HXB-2)

**Epitope** YKLKHIVWASRELER

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC DRB1\*1303, DRB1\*1502, DRB3\*0101, DRB5\*0102, DQB1\*0301, DQB1\*0601

Country United States.

Assay type CD4 T-cell Elispot - IFNγ

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 1/22 patients responded to this peptide.

**HXB2 Location** p17 (32–46)

**Author Location** p17 (32–46 B Consensus)

**Epitope** KHIVWASRELERFAV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location p17 (33-47)

Author Location p17 (33-47 IIIB, B10)

**Epitope** HIVWASRELERFAVN?

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• Peptides were identified that commonly evoke T-cell responses – 57% of 90 HIV+ people had a T-cell response to this peptide.

**HXB2 Location** p17 (33–47)

Author Location Gag (33-47 HXB-2)

**Epitope** HIVWASRELERFAVN

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** DRB1\*1303, DRB1\*1502, DRB3\*0101,

DRB5\*0102, DQB1\*0301, DQB1\*0601

Country United States.

Assay type CD4 T-cell Elispot - IFNγ

**References** Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 1/22 patients responded to this peptide.

**HXB2 Location** p17 (35–59) Author Location p17 (35-49 PV22) Epitope VWASRELERFAVNPGLLETSEGCRQ

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*13)

Donor MHC A29, A30, B8, B35, DRB1\*03, DRB1\*13

Keywords HAART, ART, Th1, Th2, TCR usage

References Lotti et al. 2002

 10/49 chronically HIV-1 infected patients had low p55-Gagspecific CD4+ T cell responses prior to therapy, and these responses remained unchanged 3 and 6 months after initiation of HAART. There was no difference in level of response in those with or without a detectable p55 response.

- For one individual, patient F45 CDC stage A2, CD4+ p55 responding clones were generated. Her response was consistently strong and heterogeneous in terms of HLA restriction and Vβ usage, and some clones had a Th1 cytokine secretion profile (high IFNgamma production) while some had a Th2 profile (high IL-4 and IL-5 production). 5/10 CD4+ clones could also induce cytotoxicity.
- 4/10 clones from patient F45 had their epitopes mapped using overlapping peptides. Clone 25 recognized this peptide sequence restricted by DRB1\*13 using TCR V $\beta$  5.1. This clone had a SI of 4.9 to p55, 13.7 to peptide, secreted low levels of IFNgamma, indicative of a Th1 response. Clone 25 had cytotoxic activity, mediated through both a perforin and a Fas-based pathway.

**HXB2 Location** p17 (37–51)

Author Location p17 (37–51 B consensus)

Epitope ASRELERFAVNPGLL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB\*0101, DRB1\*0401,

DRB1\*0405, DRB1\*0701, DRB1\*1302,

DRB1\*1501)

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This peptide was recognized by 36% of the study group.
- The peptides that were recognized by the most people were able to bind broadly to multiple HLA-DR molecules. This peptide showed high cross-reactive binding capacity and bound to 6/8 common HLA-DR molecules.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic

infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p17 (39–47)

Author Location p17 (B consensus)

**Epitope RELERFAVN** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*1302)

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI), im-

munodominance

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This epitope was in the overlap between 2 highly reactive peptides; it was fine mapped and found to be presented by DRB\*1302.

**HXB2 Location** p17 (41–51)

Author Location p17 (41-51 B consensus)

**Epitope** LERFAVNPGLL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*1302)

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This core epitope, LERFAVNPGLL, was found to bind to 1/8 HLA-DR proteins tested, DRB\*1302.

**HXB2 Location** p17 (41–55)

**Author Location** Gag (41–55 HXB-2)

Epitope LERFAVNPGLLETSE

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

DRB5\*0102, DOB1\*0301, DOB1\*0601

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ 

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 1/22 patients responded to this peptide.

**HXB2 Location** p17 (42–51)

**Author Location** p17 (B consensus)

Epitope ERFAVNPGLL

Subtype B

Immunogen HIV-1 infection

**Species (MHC)** human (DRB3\*0202, DRB3\*0301)

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI), im-

munodominance

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFN \( \gamma \) EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This epitope was in the overlap between 2 highly reactive peptides, and was fine mapped; 2 different presenting alleles for 2 different clones were determined, and found to be DRB3\*0202, DRB3\*0301.

**HXB2 Location** p17 (42–58)

**Author Location** p17 (42–58 B consensus)

Epitope ERFAVNPGLLETSEGCR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human DRB1\*0405, (DRB1\*0101,

DRB1\*1101, DRB1\*1302)

Country United States.

Assay type CD4 T-cell Elispot - IFNγ

Keywords supervised treatment interruptions (STI), im-

munodominance

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 28% of the study group.

- **Donor MHC** DRB1\*1303, DRB1\*1502, DRB3\*0101, Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
  - The peptides that were recognized by the most people were able to bind broadly to multiple HLA-DR molecules. This peptide showed high cross-reactive binding capacity and bound to 4/8 tested HLA-DR molecules.
  - The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location p17 (70-86)

**Author Location** p17 (70–86 B Consensus)

**Epitope** TGSEELRSLNTVALY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p17 (73–83)

**Author Location** Gag (73–83)

**Epitope** EELRSLYNTVA

Epitope name Gag 4.3

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT,

Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining

Keywords subtype comparisons, memory cells

References Amara et al. 2005

- Clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better-conserved for CD8 than CD4 T cells.
- 3/5 CD8 epitopes and 2/8 CD4 epitopes were conserved across multiple HIV-1 clades. All 5 CD8, and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation.
- EELRSLYNTVA was not reported for human infections.
- The response elicited to the B clade epitope EELRSLYNTVA does not cross-react with the CRF02\_AG form EEfkSLYNiVA. The epitope is however conserved across clades A,B,C,D,F.

**HXB2 Location** p17 (77–91)

Author Location Gag (77–91 HXB-2)

Epitope SLNTVATLYCVHQR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ 

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 1/22 patients responded to this peptide.

**HXB2 Location** p17 (77–94)

Author Location p17 (77-94 B consensus)

Epitope SLYNTVATLYCVHQRIEV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101, DRB1\*0401,

DRB1\*0405, DRB1\*0701, DRB1\*1302,

DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFN $\gamma$  EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This peptide was recognized by 25% of the study group.
- The peptides that were recognized by the most people were able to bind broadly to multiple HLA-DR molecules. This peptide showed high cross-reactive binding capacity and bound to 6/8 common HLA-DR molecules.

 The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location p17 (77–94)

Author Location p17 (77-94)

**Epitope** SLYNTVATLYCVHQRIEV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production

References Geels et al. 2006

- The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.
- Autologous sequences corresponding to known and predicted Th epitopes were analyzed. SLYNTVATLYCVHQRIEV had fixation of 1 mutation (SLYNT[v/i]ATLYCVHQRIEV) in 1 of the patients.

**HXB2 Location** p17 (93–107)

Author Location p17 (93–107 IIIB, B10)

Epitope EIKDTKEALDKIEEE

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location p17 (118-132)

Author Location p17 (118–132 IIIB, B10)

Epitope AAADTGHSSQVSQNY

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

## III-B-2 Gag p24 Helper/CD4+ T-cell epitopes

HXB2 Location p24 (1-9)

Author Location p24 (133-141 HXB2)

Epitope PIVQNIQGQ

**Subtype** B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101)

**Donor MHC** DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51 **Assay type** proliferation, T-cell Elispot, Intracellular cy-

tokine staining

**Keywords** HAART, ART **References** Boritz *et al.* 2003

- HIV infected individuals with advanced disease often have only weak or undetectable HIV specific Th responses, and HAART can sometimes restore Th HIV recognition in such cases. The repertoire of Th responses to p24 in an individual who responded to HAART after having a CD4+ T-cell count of 0 cells ul was determined. Eleven clonotypes were found among 13 clones, recognizing eight distinct epitopes, with a range of MHC affinities and functional avidities. Multiple Gag p24 peptides were also recognized using CD4 Elispot assays based on samples from six additional HAART-treated CD4 T-cell-reconstituted subjects.
- The TCR that recognized this epitope used  $V\beta 5.1$ .

HXB2 Location p24 (1–11)
Author Location p24 (1–11 SF2)
Epitope PIVQNLQGQMV
Immunogen HIV-1 infection
Species (MHC) human (DR1)
Keywords escape

References Harcourt et al. 1998

- 43 asymptomatic HIV+ individuals were screened for proliferative responses to HIV 12 showed a response, and dominant epitopes were mapped for two individuals, one in p24 and one in p17.
- Out of five truncated versions of peptide PIVQN-LQGQMVHQAISPRTL, only p24(1-11) elicited a proliferative response.
- Nine naturally occurring variants of this epitope were found within the individual who made this response – all bound to HLA-DR1, but three did not stimulate the CD4+ T-cell line that recognized the index peptide, suggestive of immune escape.

**HXB2 Location** p24 (1–15)

Author Location p24 (133–147 IIIB, B10)

**Epitope** PIVQNIQGQMVHQAI **Immunogen** HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

Peptides were identified that commonly evoke T-cell responses
 62% of 90 HIV+ people had a T-cell response to this peptide.

**HXB2 Location** p24 (1–22)

Author Location p24 (133-154 SF2)

Epitope PIVQNIQGQMVHQAISPRTLNA

Immunogen HIV-1 infection

Species (MHC) human

References Rosenberg et al. 1997

- While anti-HIV CD4 Th responses are characteristically undetectable in chronic infections, strong p24-specific proliferative responses were inversely correlated with low viral load in 10 chronically infected people.
- The dominant proliferative response in one of two long term survivors was to this peptide.

HXB2 Location p24 (7-21) Author Location Gag (171-185) Epitope QGQMVHQAISPRTLN

Epitope name Gag 171

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Keywords subtype comparisons

References Wilson et al. 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.
- This epitope binds to nine HLA-DR alleles: DRB1\*0101, DRB1\*1501, DRB1\*0401, DRBI\*0405, DRB1\*1302, DRB1\*0701, DRB1\*0901, DRB5\*0101 and DRB4\*0101 with an IC<sub>50</sub> threshold below 1,000 nM.
- This epitope sequence is conserved in 52% of clade B isolates.
- 7/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 non-responder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

HXB2 Location p24 (7–21) Author Location p24 (171–185)

Epitope QGQMVHQAISPRTLN

Epitope name Gag1

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Country United Kingdom.

Assay type proliferation, Intracellular cytokine staining

Keywords supertype, rate of progression

References Boaz et al. 2003

- Proliferative and cytokine (IFNgamma and II-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+ T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+ T cell declined to <500 after 15 years. Patients were treatment-naive.</li>
- Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs.
   Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.
- Gag1 was 1 of 3 peptides that had a negative correlation between absolute number of responding cells and viral load.

**HXB2 Location** p24 (7–21)

**Author Location** Gag (171–185)

Epitope QGQMVHQAISPRTLN

Epitope name Gag 171

Immunogen vaccine

mmunogen vaccine

Vector/Type: DNA with CMV promotor, peptide Adjuvant: Complete Freund's Adjuvant

(CFA)

Species (MHC) mouse (I-Ab and HLA-DR)

Donor MHC H-2b

Keywords vaccine-specific epitope characteristics, im-

munodominance

References Livingston et al. 2002

- 4 Th HIV epitopes presented by HLA-DR molecules were identified that also could be presented by murine class II molecule I-Ab, enabling testing of vaccine strategies in H-2b mice.
- Responses to pooled peptides, polyepitope peptides in a linear construct or in a branched MAP construct, and a DNA polyepitope construct with a CMV promoter were compared. A linear arrangement in polyepitope construct created a junctional epitope that could be disrupted with the addition of GPGPG spacers. The linear polyepitope construct with the GPGPG spacer worked well in terms of eliciting responses to all 4 peptides, using either DNA or protein for the vaccination.

**HXB2 Location** p24 (9-26)

**Author Location** p24 (9–26 B Consensus)

Epitope QMVHQAISPRTLNAWVKV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFN $\gamma$  EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p24 (11–26)

Author Location p24 (143–157)

Epitope VHQAISPRTLNAWVKC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

- Epitope elicits a primary proliferative response in PBMC from uninfected donors.
- Matches 3/3 anchor residues for HLA DR: VHQAISPRT

**HXB2 Location** p24 (11–30)

**Author Location** Gag (143–152 SF2)

 ${\bf Epitope} \ \ {\tt VHQAISPRTLNAWVKVVEEK}$ 

Immunogen vaccine

Vector/Type: Listeria monocytogenes Strain: B clade SF2 HIV component: p24 Gag

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Species (MHC) mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

Keywords immunodominance, Th1

References Mata & Paterson 1999
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- Listeria monocytogenes is an intracellular bacterium that lives in the cytoplasm and generates a cell-mediated immune response.
- L. monocytogenes vaccine expressing HIV-1 p24 protein (Lm-Gag) was used to stimulate gag-specific CD4+ T cell proliferative responses in BALB/c(H-2d) and C57BL/6(H-2b) mice.
- 2/3 reactive p24 peptides (out of 22 overlapping peptides that span p24) were recognized by both murine strains; this epitope is immunodominant in C57BL/6 mice and also can stimulate a BALB/c response.
- The proliferative response is due to CD4+ IFN $\gamma$ -producing cells, a Th1 response.

**HXB2 Location** p24 (11–30)

Author Location p24 (143–162 HXB2)

Epitope VHQAISPRTLNAWVKVVEEK

Subtype B

Immunogen vaccine

Vector/Type: Listeria monocytogenes Strain: B clade HXB2 HIV component: Gag

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

References Mata & Paterson 1999

- BALB/c and C57BL/6 mice were immunized with *Listeria monocytogenes* (Lm-Gag) expressing HIV-1 HXB2 Gag.
- L. monocytogenes is a gram-positive bacteria that enters the macrophage on phagocytosis and lives in the cytoplasm; secreted *L. monocytogenes* antigens are processed and presented by both class I and class II pathways.
- The class II Th response was probed using 20mer peptides that overlapped by 10; the peptides VHQAISPRTL-NAWVKVVEEK and FRDYVDRFYKTLRAEQASQD were recognized in H-2<sup>b</sup> and H-2<sup>d</sup> mice.

**HXB2 Location** p24 (21–36)

Author Location p24 (153–167)

Epitope NAWVKVVEEKAFSPEC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

• Epitope elicits a primary proliferative response in PBMC from uninfected donors.

**HXB2 Location** p24 (23–40)

**Author Location** p24 (23–40 B Consensus)

Epitope WVKVVEEKAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p24 (23–40) **Author Location** p24 (23–40)

Epitope WKVVEEKAFSPEVIPMF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production

References Geels et al. 2006

- The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.
- Autologous sequences corresponding to known and predicted Th epitopes were analyzed. WKVVEEKAFSPEVIPMF had fixation of 2 mutations WKV[v/i]EEKAF[s/n]PEVIPMF in 1 of the patients.

HXB2 Location p24 (25-39)

**Author Location** 

Epitope KVVEEKAFSPEVIPM

Epitope name G040

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine

assay

Keywords memory cells

References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-γ, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

**HXB2 Location** p24 (28–36)

Author Location p24 (160–168 HXB2)

**Epitope** EEKAFSPEV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101)

**Donor MHC** DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51 **Assay type** proliferation, T-cell Elispot, Intracellular cy-

tokine staining

Keywords HAART, ART

References Boritz et al. 2003

- HIV infected individuals with advanced disease often have only weak or undetectable HIV specific Th responses, and HAART can sometimes restore Th HIV recognition in such cases. The repertoire of Th responses to p24 in an individual who responded to HAART after having a CD4+ T-cell count of 0 cells ul was determined. Eleven clonotypes were found among 13 clones, recognizing eight distinct epitopes, with a range of MHC affinities and functional avidities. Multiple Gag p24 peptides were also recognized using CD4 Elispot assays based on samples from six additional HAART-treated CD4 T-cell-reconstituted subjects.
- The TCR that recognized this epitope used  $V\beta 2$ .

**HXB2 Location** p24 (28–38)

**Author Location** p24 (HXB2)

**Epitope** EEKAFSPEVIP

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DQ5)

Assay type CD4 T-cell Elispot - IFN $\gamma$ 

Keywords epitope processing, vaccine antigen design

References SenGupta et al. 2004

Multiple HLA calss I-restricted and class II-restricted T-cell
epitopes were shown to be processed and presented from an
exogenously added HIV-1 gag-p24 peptide complexed to a
heat shock protein. T-cell recognition of the complex was
shown to be inhibited by brefeldin A indicating an endoplasmic
reticulum-dependent pathway.

**HXB2 Location** p24 (28–38)

**Author Location** p24 (161–171 NY-5)

Epitope EEKAFSPEVIP

Epitope name EP11

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DQ5)

Donor MHC DR11, DR14, Drw52, DQ3, DQ5

Country United States.

Assay type proliferation, CD4 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

Keywords subtype comparisons, rate of progression,

acute/early infection, early treatment, variant cross-recognition or cross-neutralization

References Norris et al. 2004

• 5 CD4-T cell epitopes in Gag-p24 were studied, and a minimum epitope length of 6-16 amino acids was found to be required for CD4+ T cell proliferation. Cross-clade recognition was found to be impaired in 17/32 variants tested.

- Patient AC-01, who was infected with HIV-1 in 1997, recognized this epitope and epitope EPRGSDIAGT during acute infection, and 19 months post-initiation of ART therapy started during primary infection.
- The epitope EEKAFSPEVIP is highly conserved in B clade. Common variants from other clades were tested and all had markedly diminished responses, including eeRafspevip, eekaLspevip, and eDkafspevip (all found in clade A); eekGfspevip, eekGfNpevip (clades A and CRF01\_AE); eekafspeIip (clade C); eekafNpevip (clade D).
- Minimum length peptides for the epitopes studied were not particularly optimal, and peptides longer than the minimum, up to 22 amino acids, were often as potent, in marked contrast to CD8+ T-cell responses.

HXB2 Location p24 (31-46) Author Location p24 (163–177) Epitope AFSPEVIPMFSALSEC

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201) References Bedford et al. 1997

- Epitope elicits a primary proliferative response in PBMC from uninfected donors.
- Peptide contains a CTL epitope identified in HIV-positive pa-
- Peptide binds to HLA A\*0201 and causes regulation of class I expression on T2 cells.
- Matches 3/3 anchor residues for HLA DR: VIPMFSALS

**HXB2 Location** p24 (31–47)

**Author Location** p24 (31–47 B Consensus)

Epitope AFSPEVIPMFSALSEGA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNy EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p24 (31–48)

Author Location p24 (31-48)

Epitope AFSPEVIPMFSALSEGAT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production

References Geels et al. 2006

- The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.
- · Autologous sequences corresponding to known and predicted Th epitopes were analyzed. AFSPEVIPMFSALSEGAT had fixation of 2 mutations (AF[s/n]PEVIPMF[s/t]ALSEGAT) in 1 of the patients.

**HXB2 Location** p24 (31–52)

Author Location p24 (163–184 SF2)

Epitope AFSPEVIPMFSALSEGATPQDL

Immunogen HIV-1 infection

Species (MHC) human

References Rosenberg et al. 1997

- · Low viral load correlated with strong HIV-1-specific proliferative response.
- · A proliferative response to this epitope was detected in two long term survivors.

**HXB2 Location** p24 (34–49)

**Author Location** p24 (HXB2)

**Epitope** PEVIPMFSALSEGATP

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DR1)

Assav type CD4 T-cell Elispot - IFNγ

Keywords epitope processing, vaccine antigen design,

characterizing CD8+ T cells

References SenGupta et al. 2004

• Multiple HLA class I-restricted and class II-restricted T-cell epitopes were shown to be processed and presented from an exogenously added HIV-1 gag-p24 peptide complexed to a heat shock protein. T-cell recognition of the complex was shown to be inhibited by brefeldin A indicating an endoplasmic reticulum-dependent pathway.

**HXB2 Location** p24 (34–49)

**Author Location** p24 (168–177 NY-5)

Epitope PEVIPMFSALSEGATP

Epitope name PP16

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DR1)

Donor MHC DR1, DR11, DRw52, DQ5, DQ7

Country United States.

Assay type proliferation, CD4 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

**Keywords** rate of progression, acute/early infection, early treatment, variant cross-recognition or cross-neutralization

References Norris et al. 2004

- 5 CD4-T cell epitopes in Gag-p24 were studied, and a minimum epitope length of 6-16 amino acids was found to be required for CD4+ T-cell proliferation. Cross-clade recognition was studied and found to be impaired in 17/32 variants tested.
- Patient AC-25 was an acute seroconvertor at the time of sampling, infected with HIV-1 in 1998, and given ARVs during primary infection. The study subject was resampled 18 months after initiation of therapy.
- · Natural variants of the epitope PEVIPMFSALSEGATP diminished the level of the response, including pevipmfPalsegStp and pevipmfsalsegStp, found in CRF01\_AE; peIipmfTalsegatp, clade C; pevipmfsalSegatp, clade B; pevipmfTalsegatp, clades A, B and C; and pevipVfsalsegatp, clade A.
- Minimum length peptides for the epitopes studies were not particularly optimal, and peptides longer than the minimum, up to 22 amino acids, were often as potent, in marked contrast to CD8+ T-cell responses.

HXB2 Location p24 (34-49)

Author Location p24

Epitope PEVIPMFSALSEGATP

Epitope name PP16

Immunogen HIV-1 infection Species (MHC) human (DR1)

> Elispot - IFNγ, Flow cytometric T-cell cytokine assay

Keywords binding affinity, escape, optimal epitope References Norris et al. 2006

- This study demonstrates a mechanism of antagonism by a peptide shorter than the minimum length epitope for an HIV p24-specific CD4+ T-cell clone.
- Truncation of the peptide from PEVIPMFSALSEGATP (PP16) to PEVIPMFSALSEG (PG13) rendered the peptide unable to elicit proliferation, IFNy release, or serine esterase release, even though it retained strong binding to MHC.
- Although both the original and truncated peptide-MHC complexes bound TCR clone, PEVIPMFSALSEGATP-DR1 tetramer bound with higher avidity than PEVIPMFSALSEG-DR1 tetramer, suggesting that tighter association of the peptide-MHC complex with the TCR is associated with the extent of T-cell activation.
- G/P substitution in the original full-length peptide (PEVIPMF-SALSE[g/p]ATP) led to complete loss of agonist activity, and PEVIPMFSALSEpATP became antagonistic.

HXB2 Location p24 (35-44)

**Author Location** p24 (HXB2)

**Epitope** EVIPMFSALS

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DR4)

Assay type CD4 T-cell Elispot - IFNγ

Keywords epitope processing, vaccine antigen design,

characterizing CD8+ T cells

References SenGupta et al. 2004

• Multiple HLA calss I-restricted and class II-restricted T-cell epitopes were shown to be processed and presented from an exogenously added HIV-1 gag-p24 peptide complexed to a heat shock protein. T-cell recognition of the complex was shown to be inhibited by brefeldin A indicating an endoplasmic reticulum-dependent pathway.

**HXB2 Location** p24 (35–44)

Author Location p24 (168-177 NY-5)

**Epitope** EVIPMFSALS

Epitope name ES10

Subtype B

Immunogen HIV-1 infection Species (MHC) human (DR4)

Donor MHC DR4, DR15, DRw51, DRw53, DQ3, DQ6

Country United States.

Assay type proliferation, CD4 T-cell Elispot - IFNγ,

Chromium-release assay

Keywords rate of progression, acute/early infection, early treatment, variant cross-recognition or

cross-neutralization

References Norris et al. 2004

- 5 CD4-T cell epitopes in Gag-p24 were studied, and a minimum epitope length of 6-16 amino acids was found to be required for CD4+ T-cell proliferation. Cross-clade recognition was studied and found to be impaired in 17/32 variants tested.
- Patient 161J, infected with HIV-1 in the mid 1980s, was 1 of the 2 LTNPs examined. 161J was ART naive.
- Assay type proliferation, Tetramer binding, CD4 T-cell Natural variants of the epitope EVIPMFSALS gave diminished responses including evipmfTals, common in clades A, B and C; and evipVfsals, clade A; evipmfsalA, a clade B variant; and elipmfTals, clade C. The exception was the CRF01\_AE variant evipmfPals, which was as reactive as the original peptide tested.
  - Minimum length peptides for the epitopes studies were not particularly optimal, and peptides longer than the minimum, up to 22 amino acids, were often as potent, in marked contrast to CD8+ T-cell responses.

**HXB2 Location** p24 (35–44)

Author Location p24 (167–176 HXB2)

**Epitope** EVIPMFSALS

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101)

Donor MHC DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51

Assay type proliferation, T-cell Elispot, Intracellular cy-

tokine staining

Keywords HAART, ART

References Boritz et al. 2003

· HIV infected individuals with advanced disease often have only weak or undetectable HIV specific Th responses, and HAART can sometimes restore Th HIV recognition in such cases. The repertoire of Th responses to p24 in an individual who responded to HAART after having a CD4+ T-cell count of 0 cells ul was determined. Eleven clonotypes were found among 13 clones, recognizing eight distinct epitopes, with a range of MHC affinities and functional avidities. Multiple Gag p24 peptides were also recognized using CD4 Elispot assays based on samples from six additional HAART-treated CD4 T-cell-reconstituted subjects.

HXB2 Location p24 (41–56)

Author Location p24 (173–187)

Epitope SALSEGATPQDLNTMC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

 Epitope elicits a primary proliferative response in PBMC from uninfected donors.

HXB2 Location p24 (48–62)
Author Location p24 (180–194)
Epitope TPQDLNTMLNTVGGH
Immunogen HIV-1 infection
Species (MHC) human

References Adams et al. 1997

- One of four immunogenic Gag peptides used in study of proliferative response to p24.
- Homology to an SIV epitope recognized by macaque T-cells.
- T-cells from 8 of 19 HIV+ individuals responded to this epitope.
- Improved assay system (increase in culture time to 8 days and addition of IL-2 to cultures) gave increased detection of proliferative response.

HXB2 Location p24 (51–66)
Author Location p24 (183–197)
Epitope DLNTMLNTYGGHQAAC
Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

 Epitope elicits a primary proliferative response in PBMC from uninfected donors.

**HXB2 Location** p24 (51–82) **Author Location** Gag (183–214 LAI)

Epitope DLNTMLNTVGGHQAAMQMLKETINEEAAEWDR

Subtype B Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I trial.
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees 2/10 reacted to this peptide.
- 9/12 tested mounted a CTL responses to at least one of the six peptides, each of the six peptides elicited a CTL response in at least one individual.
- None of the 12 tested had an IgG response to this peptide.

HXB2 Location p24 (69–88)

Author Location p24 (201–220 IIIB)

Epitope LKETINEEAAEWDRVHPVHA

Epitope name P21

Immunogen in vitro stimulation or selection

Species (MHC) human (DR)

Donor MHC DR4, DR7, DQ2, DQ3

Keywords immunodominance, Th1, Th2, TCR usage

References Venturini et al. 2002

- PBMC from a seronegative donor, the healthy brother of a pair
  of monozygotic twins discordant for HIV-1 infection, were
  used to generate HIV-1 Gag-specific CD4+ T-cell clones by
  in vitro immunization with HIV-1 overlapping 20mer peptides
  spanning p55. Six clones were generated by limiting dilution.
  All reacted with p24 except one which recognized a p24 peptide
  and a p6 peptide. All CD4+ T cell clones were HLA class II
  DR restricted.
- Clone 85 recognized this peptide using TCR V $\beta$  8 and 18; the two TCR receptors indicates this limiting dilution represents a mixed population. This clone was only activated by peptide, not by processed protein from vaccinia virus Gag-infected B-LCL.

**HXB2 Location** p24 (71–86) **Author Location** p24 (203–220)

Epitope ETINEEAAEWDRVHPC

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

• Epitope elicits a primary proliferative response in PBMC from uninfected donors.

HXB2 Location p24 (71-88)

Author Location p24

Epitope ETINEEAAEWDRVHPVHA

Epitope name 17 Subtype B Immunogen

Species (MHC) (DRB1\*0101)

**Donor MHC** DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51

References

**HXB2 Location** p24 (71–88) **Author Location** (203–220)

Epitope ETINEEAAEWDRVHPVHA

Subtype B Immunogen

Species (MHC) human (DRB1\*0101)

Donor MHC DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51

References

**HXB2 Location** p24 (71–88)

**Author Location** 

Epitope ETINEEAAEWDRVHPVHA

Epitope name 17
Subtype B
Immunogen

Species (MHC) (DRB1\*0101)

Donor MHC DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51

References

**HXB2 Location** p24 (71–88)

Author Location p24 (203–220 HXB2)

Epitope ETINEEAAEWDRVHPVHA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101)

Donor MHC DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51

tokine staining

Keywords HAART, ART References Boritz et al. 2003

- · HIV infected individuals with advanced disease often have only weak or undetectable HIV specific Th responses, and HAART can sometimes restore Th HIV recognition in such cases. The repertoire of Th responses to p24 in an individual who responded to HAART after having a CD4+ T-cell count of 0 cells/ul was determined. Eleven clonotypes were found among 13 clones, recognizing eight distinct epitopes, with a range of MHC affinities and functional avidities. Multiple Gag p24 peptides were recognized using CD4 Elispot assays based on samples from six additional HAART-treated CD4 T-cell-reconstituted subjects.
- The Th clone that recognized this epitope utilized TCR V $\beta$ 17.

HXB2 Location p24 (71-92)

Author Location p24 (203–224 HXB2)

Epitope ETINEEAAEWDRVHPVHAGPIA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101)

**Donor MHC** DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51

Assay type proliferation, T-cell Elispot, Intracellular cy-

tokine staining Keywords HAART, ART

References Boritz et al. 2003

• HIV infected individuals with advanced disease often have only weak or undetectable HIV specific Th responses, and HAART can sometimes restore Th HIV recognition in such cases. The repertoire of Th responses to p24 in an individual who responded to HAART after having a CD4+ T-cell count of 0 cells/ul was determined. Eleven clonotypes were found among 13 clones, recognizing eight distinct epitopes, with a range of MHC affinities and functional avidities. Multiple Gag p24 peptides were recognized using CD4 Elispot assays based on samples from six additional HAART-treated CD4 T-cell-reconstituted subjects.

**HXB2 Location** p24 (73–83) Author Location Gag (205-215)

**Epitope INEEAAEWDRV** 

**Epitope name** Gag 11.2

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT,

Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining

**Keywords** subtype comparisons, memory cells

References Amara et al. 2005

• Clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better-conserved for CD8 than CD4 T cells.

- Assay type proliferation, T-cell Elispot, Intracellular cy- 3/5 CD8 epitopes and 2/8 CD4 epitopes were conserved across multiple HIV-1 clades. All 5 CD8, and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation.
  - CD4 T-cell epitope previously reported for human is ETINEEAAEWDRVHPC. HLA restriction: DRB1\*03, DRB1\*0101.
  - The response elicited to the B clade epitope INEEAAEWDRV does not cross-react with the CRF02\_AG form INdEAAEW-DRV. The epitope is however conserved in CRF01\_AE and CRF02\_AG consensus sequences. Other clades tend to have L in the last position (INEEAAEWDR[v/l]).

**HXB2 Location** p24 (73–97)

**Author Location** p24 (205–229 PV22)

Epitope INEEAAEWDRVHPVHAGPIAPGQMR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*03)

Donor MHC A29, A30, B8, B35, DRB1\*03, DRB1\*13

Keywords HAART, ART, Th1, Th2, TCR usage

References Lotti et al. 2002

- 10/49 chronically HIV-1 infected patients had low p55-Gagspecific CD4+ T cell responses prior to therapy, and these responses remained unchanged 3 and 6 months after initiation of HAART. There was no difference in level of response in those with or without a detectable p55 response.
- For one individual, patient F45 CDC stage A2, CD4+ p55 responding clones were generated. Her response was consistently strong and heterogeneous in terms of HLA restriction and  $V\beta$ usage, and some clones had a Th1 cytokine secretion profile (high IFNgamma production) while some had a Th2 profile (high IL-4 and IL-5 production). 5/10 CD4+ clones could also induce cytotoxicity.
- 4/10 clones from patient F45 had their epitopes mapped using overlapping peptides. Clone 12 recognized this peptide sequence restricted by DRB1\*03 using TCR Vβ 22. This clone had a SI of 12.4 to p55, 49.6 to peptide, secreted low levels of IFNgamma, indicative of a Th1 response. Clone 12 had cytotoxic activity, mediated through both a perforin and a Fas-based pathway.

**HXB2 Location** p24 (76–85)

Author Location p24 (208–217)

**Epitope** EAAEWDRVHP

Immunogen HIV-1 infection

Species (MHC) human

References Adams et al. 1997

- One of four immunogenic Gag peptides used in study of the proliferative response to p24.
- T-cells from 11 of 24 HIV+ individuals responded to this epi-
- Improved assay system (increase in culture time to 8 days and addition of IL-2 to cultures) gave increased detection of proliferative response.

**HXB2 Location** p24 (76–90)

Author Location p24 (208–222 IIIB, B10)

Epitope EAAEWDRVHPVHAGP

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location p24 (79-88)

Author Location p24 (211–220 HXB2)

Epitope EWDRVHPVHA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101)

**Donor MHC** DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51 Assay type proliferation, T-cell Elispot, Intracellular cy-

tokine staining

Keywords HAART, ART References Boritz et al. 2003

· HIV infected individuals with advanced disease often have only weak or undetectable HIV specific Th responses, and HAART can sometimes restore Th HIV recognition in such cases. The repertoire of Th responses to p24 in an individual who responded to HAART after having a CD4+ T-cell count of 0 cells ul was determined. Eleven clonotypes were found among 13 clones, recognizing eight distinct epitopes, with a range of MHC affinities and functional avidities. Multiple Gag p24 peptides were also recognized using CD4 Elispot assays based on samples from six additional HAART-treated CD4 T-cell-reconstituted subjects. Two clones recognized this epitope.

**HXB2 Location** p24 (81–95)

Author Location p24 (215-229 SF2)

Epitope DRVHPVHAGPIAPGQ

Immunogen vaccine

*Vector/Type:* virus-like particle (VLP) Strain: B clade SF2 HIV component: p24 Gag

Species (MHC) macaque

References Mills et al. 1990

• Responses to 3 T-cell and multiple linear B-cell epitopes were found in vaccinated macaques.

**HXB2 Location** p24 (81–102)

Author Location p24 (213–234 SF2)

Epitope DRVHPVHAGPIAPGQMREPRGS

Immunogen HIV-1 infection

Species (MHC) human

References Rosenberg et al. 1997

- · While anti-HIV CD4 Th responses are characteristically undetectable in chronic infections, strong p24-specific proliferative responses were inversely correlated with low viral load in 10 chronically infected people.
- The dominant proliferative response in one of two long term survivors was to this peptide.

HXB2 Location p24 (81–102)

Author Location p24 (81–102)

Epitope DRVHPVHAGPIAVPGQMREPRGS

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production

References Geels et al. 2006

- The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.
- Autologous sequences corresponding to known and predicted Th epitopes were analyzed. DRVHPVHAGPI-AVPGQMREPRGS had fixation of 1 mutation (DRVH-PVHAGPI[a/p]VPGQMREPRGS) in 1 of the patients.

**HXB2 Location** p24 (85–99)

**Author Location** 

Epitope PVHGPIAPGQMREP

Epitope name G055

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine assay

Keywords memory cells

References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-y, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

HXB2 Location p24 (86–94)

Author Location p24 (NY5)

Epitope VHAGPIAPG

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DO7)

Keywords HAART, ART, supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection, crosspresentation by different HLA, early treatment, TCR usage

References Norris et al. 2001

• Gag-specific CD4+ helper T-cell clones were derived from 1 LTNP (CTS-01) and 3 individuals given therapy during acute infection, 2 before (AC-01 and AC-36) and 1 after (AC-25) STI. Gag peptide recognition induced proliferation, IFNγ production, and perforin-mediated cytotoxicity in all CD4+ T-cell clones isolated.

- 3/23 p24-derived peptides tested induced proliferative p24- HIV-1-specific CD4+ T-cell responses were analyzed for 6 specific Th cell responses in the LTNP CTS-01. The immunodominant response was to the peptide DRVHPVHAGPI-APGQMREPRGS (81-102), and 9/10 CD4+ T-cell clones reacted with it. One was characterized in detail and used a B $\beta$ 4 TCR.
- The minimum peptide recognized by the clones from CTS-01 was VHAGPIAPG and it was restricted by HLA-DQ7.

**HXB2 Location** p24 (86–94)

**Author Location** p24 (219–227 NY-5)

Epitope VHAGPIAPG

Epitope name VG9

Subtype B

Immunogen HIV-1 infection Species (MHC) human (DQ7)

Donor MHC DR11, DR15, DRw51, DRw52, DQ6, DQ7

Country United States.

Assay type proliferation, CD4 T-cell Elispot - IFNγ Keywords acute/early infection, early treatment, variant cross-recognition or cross-neutralization

References Norris et al. 2004

- 5 CD4-T cell epitopes in Gag-p24 were studied, and a minimum epitope length of 6-16 amino acids was found to be required for CD4+ T-cell proliferation. Cross-clade recognition was studied and found to be impaired in 17/32 variants tested.
- Patient CTS01, who was infected with HIV-1 in 1998, was an LTNP, and recognized this epitope.
- This epitope VHAGPIAPG was the most variable of the 5 epitopes studied. Only the C variant Ihagpiapg did not diminish the response. All other variations had impaired responses: vhagpVapg, found in clades A, B, C, and D; vQagpVapg, clades B, C, D; AQagpFPpg, IhagpVapg, AhagpVapg, and vQagpiP, all found in clade A; AQagpiapg, clade B; and vPagpiapg, clade C.
- Minimum length peptides for the epitopes studies were not particularly optimal, and peptides longer than the minimum, up to 22 amino acids, were often as potent, in marked contrast to CD8+ T-cell responses.

HXB2 Location p24 (87-101)

Author Location p24 (219–233 BRU)

Epitope HAGPIAPGQMREPRG

Immunogen in vitro stimulation or selection

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Vaslin et al. 1994

• Peptide G2: could prime for in vitro immunoproliferative responses and for subsequent IgG responses.

HXB2 Location p24 (93-107)

**Author Location** 

Epitope PGQMREPRGSDIAGT

Epitope name G057

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine

assay

Keywords memory cells References Younes et al. 2003

- years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-γ, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

**HXB2 Location** p24 (96–103)

Author Location p24 (228–235 LAI)

Epitope MREPRGSD

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

**HXB2 Location** p24 (96–110)

Author Location p24 (228–242 IIIB, B10)

**Epitope** MREPRGSKIAGTTST

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location p24 (98–107)

**Author Location** p24 (231–240 NY-5)

Epitope EPRGSDIAGT

Epitope name ET10

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DQ7)

Donor MHC DR11, DR14, Drw52, DQ3, DQ5

Country United States.

Assay type proliferation, CD4 T-cell Elispot - IFN $\gamma$ ,

Chromium-release assay

Keywords acute/early infection, early treatment, variant cross-recognition or cross-neutralization

References Norris et al. 2004

- 5 CD4-T cell epitopes in Gag-p24 were studied, and a minimum epitope length of 6-16 amino acids was found to be required for CD4+ T-cell proliferation. Cross-clade recognition was found to be impaired in 17/32 variants tested.
- Patient AC-01, who was infected with HIV-1 in 1997, recognized this epitope and epitope EPRGSDIAGT during acute infection, and 19 months post-initiation of ART therapy started during primary infection.
- This was the most variable of the 5 epitopes studied. REPRGS-DIAGT natural variants were tested and did not usually diminish the response by much (rDprgsdiagt, clades B and C; and rDprgsdiagA and rGprgsdiagt, both clade C), although the CRF01\_AE variant reprgAdiagt abrogated the response.

• Minimum length peptides for the epitopes studies were not particularly optimal, and peptides longer than the minimum, up to 22 amino acids, were often as potent, in marked contrast to CD8+ T cell responses. This peptide was the exception, as REPRGSDIAGTT, which is elongated by 2 amino acids compared to the minimum epitope, elicited a stronger proliferative immune response as well as IFN $\gamma$  secretion and cytolyis.

HXB2 Location p24 (99-118)

Author Location p24 (231-250 IIIB)

Epitope PRGSDIAGTTSTLQEQIGWM

Epitope name P24

Immunogen in vitro stimulation or selection

Species (MHC) human (DR4)

Donor MHC DR4, DR7, DQ2, DQ3

Keywords immunodominance, Th1, Th2, TCR usage

References Venturini et al. 2002

- PBMC from a seronegative donor, the healthy brother of a pair of monozygotic twins discordant for HIV-1 infection, were used to generate HIV-1 Gag-specific CD4+ T-cell clones by in vitro immunization with HIV-1 overlapping 20mer peptides spanning p55. Six clones were generated by limiting dilution. All reacted with p24 except one which recognized a p24 peptide and a p6 peptide. All CD4+ T cell clones were HLA class II DR restricted.
- Clone 6 recognized three peptides including this one with a Th1 response using TCR V $\beta$  6 (6s5A1N1). Sequencing TCR V $\beta$ regions of colonies from clone 6 suggested this was a clonal population. Assays using different peptide concentrations suggest that this peptide does not carry the main epitope of this clone.

HXB2 Location p24 (101-115)

Author Location p24 (235–249 SF2)

Epitope GSDIAGTTSTLQEQI

Immunogen vaccine

Vector/Type: virus-like particle (VLP) Strain: B clade SF2 HIV component: p24 Gag

Species (MHC) macaque

References Mills et al. 1990

• Responses to 3 T-cell and multiple linear B-cell epitopes were found in vaccinated macaques - epitope response defined by T-cell clone.

**HXB2 Location** p24 (101–115)

**Author Location** Gag (233–247 HXB-2)

Epitope GSDIAGTTSTQEQI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC DRB1\*1303, DRB1\*1502, DRB3\*0101,

DRB5\*0102, DQB1\*0301, DQB1\*0601

Country United States.

Assay type CD4 T-cell Elispot - IFNγ References Koeppe et al. 2006

• The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.

1/22 patients responded to this peptide.

HXB2 Location p24 (101–116)

Author Location p24

Epitope GSDIAGTTSTLQEQIC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

• Epitope elicits a primary proliferative response in PBMC from uninfected donors.

**HXB2 Location** p24 (109–123)

**Author Location** 

Epitope STLQEQIGWMTNNPP

Epitope name G061

Immunogen HIV-1 infection

Species (MHC) human Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine

assay

Keywords memory cells References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN- $\gamma$ , but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

HXB2 Location p24 (109–128)

Author Location p24 (241–260 IIIB)

Epitope STLQEQIGWMTNNPPIPVGE

Epitope name P25

Immunogen in vitro stimulation or selection

Species (MHC) human

Donor MHC DR4, DR7, DQ2, DQ3

Keywords immunodominance, Th1, Th2, TCR usage

References Venturini et al. 2002

• PBMC from a seronegative donor, the healthy brother of a pair of monozygotic twins discordant for HIV-1 infection, were used to generate HIV-1 Gag-specific CD4+ T-cell clones by in vitro immunization with HIV-1 overlapping 20mer peptides spanning p55. Six clones were generated by limiting dilution. All reacted with p24 except one which recognized a p24 peptide and a p6 peptide. All CD4+ T cell clones were HLA class II DR restricted.

• Clone 50 recognized this peptide with a Th0 response (Th0 means that cytokines characteristic of both Th1 and Th2 responses were stimulated), using TCR V $\beta$  17, and was a homogeneous T-cell population. This clone was only activated by peptide, not by processed protein from vaccinia virus Gaginfected B-LCL.

**HXB2 Location** p24 (111–132) **Author Location** p24 (243–264 SF2)

Epitope LQEQIGWMTNNPPIPVGEIYKR

Immunogen HIV-1 infection

Species (MHC) human

References Rosenberg et al. 1997

- Low viral load correlated with strong HIV-1-specific proliferative response.
- A proliferative response to this epitope was detected in two long term survivors.

HXB2 Location p24 (119–133)

Author Location p24 (251–265)

Epitope TNNPPIPBGEIYKRW

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*1301)

Konwords binding offinity HAA

Keywords binding affinity, HAART, ART

**References** Blankson & Siliciano 2001; Malhotra *et al.* 2001

- The DRB1\*13-DQB1\*06 haplotype is associated with maintained viral suppression after HAART 7/7 early-treated DRB1\*13-DQB1\*06 positive people, but only 3/14 (21%) of those who did not have DRB1\*13-DQB1\*06, maintained viral suppression for 18 months.
- PBMC from individuals with the haplotype DRB1\*13-DQB1\*06 displayed increased IFNγ secretion and stronger proliferative responses against p24 80 weeks post treatment.
- DRB1\*13-DQB1\*06 was also found to be enriched among long-term non-progressors (LTNPs) (it was in 9/18 versus, versus 21% of the general population)
- This epitope was mapped with truncated peptides using the Elispot assay.
- Two distinct DRB1\*13 epitopes were defined in the peptide region spanning 251 to 270, and this 20-mer bound with very high affinity to DRB1\*1302 DRB1\*1301 and DRB1\*1302 would be expected to have very similar binding properties.

**HXB2 Location** p24 (119–133) **Author Location** p24 (119–133)

Epitope TNNPPIPBGEIYKRW

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production

References Geels et al. 2006

The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.

 Autologous sequences corresponding to known and predicted T-helper epitopes were analyzed. TNNPPIPBGEIYKRW had fixation of R14K mutation (TNNPPIPBGEIYKkW) in 1 of the patients.

**HXB2 Location** p24 (121–136)

**Author Location** p24 (253–267)

Epitope NPPIPVGEIYKRWIIC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

 Epitope elicits a primary proliferative response in PBMC from uninfected donors.

**HXB2 Location** p24 (121–140)

Author Location Gag (253–272 SF2)

Epitope NPPIPVGEIYKRWILGLNK

Immunogen vaccine

Vector/Type: Listeria monocytogenes Strain: B clade SF2 HIV component: p24 Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

**Keywords** immunodominance, Th1 **References** Mata & Paterson 1999

- Listeria monocytogenes is an intracellular bacterium that lives in the cytoplasm and generates a cell-mediated immune response.
- *L. monocytogenes* vaccine expressing HIV-1 p24 protein (Lm-Gag) was used to stimulate gag specific CD4+ T cell proliferative responses in BALB/c(H-2d) and C57BL/6(H-2b) mice.
- 2/3 reactive p24 peptides (out of 22 overlapping peptides that span p24) were recognized by both murine strains; this epitope is immunodominant in BALB/c mice and did not stimulate a C57BL/6 response.
- The proliferative response is due to CD4+ IFN $\gamma$ -producing cells, a Th1 response.

**HXB2 Location** p24 (121–140)

Author Location p24 (253–272 HXB2)

Epitope NPPIPVGEIYKRWIILGLNK

Subtype B

Immunogen vaccine

Vector/Type: Listeria monocytogenes Strain: B clade HXB2 HIV component: Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords immunodominance

References Mata & Paterson 1999

- BALB/c and C57BL/6 mice were immunized with *Listeria monocytogenes* (Lm-Gag) expressing HIV-1 HXB2 Gag.
- L. monocytogenes is a gram-positive bacteria that enters the macrophage on phagocytosis and lives in the cytoplasm; secreted *L. monocytogenes* antigens are processed and presented by both class I and class II pathways.
- The class II Th response was probed using 20mer peptides that overlapped by 10; the peptide MPPIPVGEIYKRWIILGLNK gave the immunodominant response for the H-2<sup>d</sup> haplotype, but was not recognized in H-2<sup>b</sup> mice.

**HXB2 Location** p24 (121–140)

Author Location Gag (197-205)

Epitope NPPIPVGEIYKRWIILGLNK

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade

HXB2 HIV component: Gag

**Species (MHC)** mouse (H-2d) **Country** United States.

Assay type proliferation, T-cell Elispot

**Keywords** vaccine antigen design

References Kwak et al. 2004

 A recombinant vaccinia virus with HIV-1 Gag replacing the cytoplasmic domain of the B5R protein was shown to induce better primary CD4 response than recombinant vaccinia virus expressing Gag from the TK-locus; CD8 responses were less specific. When immunized BALB/c mice were challenged with a recombinant Listeria that expresses HIV-Gag, lower colony counts of Listeria were found in the liver and spleen of mice immunized with virus expressing B5R-Gag fusion protein.

**HXB2 Location** p24 (121–152)

Author Location Gag (183–214 LAI)

Epitope NPPIPVGEIYKRWIILGLNKIVRMYSPTSILD

Subtype B Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I trial.
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees 9/10 reacted to this peptide.
- 9/12 tested mounted a CTL responses to at least one of the six peptides, each of the six peptides elicited a CTL response in at least one individual – this peptide was particularly immunogenic, eliciting a CTL response in four vaccinees.
- All of the 12 tested had an IgG response to this peptide.

**HXB2 Location** p24 (125–139)

**Author Location** Gag (257–271 HXB-2)

Epitope PVGEIYKRWIILGLN

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** DRB1\*1303, DRB1\*1502, DRB3\*0101,

DRB5\*0102, DQB1\*0301, DQB1\*0601;

Country United States.

Assay type CD4 T-cell Elispot - IFNγ

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 2/22 patients responded to this peptide.

**HXB2 Location** p24 (125–139)

**Author Location** 

Epitope PVGEIYKRWIILGLN

Epitope name G065

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

**Assay type** proliferation, Flow cytometric T-cell cytokine

assay

Keywords memory cells

References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-γ, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

**HXB2 Location** p24 (127–141)

Author Location Gag (294–308)

Epitope GEIYKRWIILGLNKI

Epitope name Gag 294

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Keywords subtype comparisons

References Wilson et al. 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicted proliferative responses from multiple HIV-infected donors.
- This epitope binds ten HLA-DR alleles: DRB1\*0101, DRB1\*1501, DRB1\*0405, DRB1\*1101, DRB1\*1302, DRB1\*0701, DRB1\*0802, DRB1\*0901, DRB5\*0101 and DRB4\*0101 with an IC<sub>50</sub> threshold below 1,000 nM.
- This epitope sequence is conserved in 95% of clade B isolates.
- 6/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 non-responder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

**HXB2 Location** p24 (127–141)

**Author Location** p24 (294–308)

Epitope GEIYKRWIILGLNKI

Epitope name Gag2

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Country United Kingdom.

Assay type Cytokine production, proliferation

Keywords supertype, rate of progression

References Boaz et al. 2003

Proliferative and cytokine (IFNgamma and Il-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+ T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+

T cell declined to <500 after 15 years. Patients were treatment-naive.

Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs.
 Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.

HXB2 Location p24 (128–137) Author Location p24 (260–269) Epitope EIYKRWIILG Immunogen HIV-1 infection

**Species (MHC)** human (DRB1\*1301, DRB1\*1302)

Keywords binding affinity, HAART, ART, Th1

**References** Blankson & Siliciano 2001; Malhotra *et al.* 2001

- The DRB1\*13-DQB1\*06 haplotype is associated with maintained viral suppression after HAART 7/7 early-treated DRB1\*13-DQB1\*06 positive people, but only 3/14 (21%) of those who did not have DRB1\*13-DQB1\*06, maintained viral suppression for 18 months.
- PBMC from individuals with the haplotype DRB1\*13-DQB1\*06 displayed increased IFNγ secretion and stronger proliferative responses against p24 80 weeks post treatment.
- DRB1\*13-DQB1\*06 was also found to be enriched among long-term non-progressors (it was in 9/18 versus, versus 21% of the general population)
- The truncated peptide that gave the optimal proliferative response for a Th1 phenotype clone was this nine-mer.
- This region, shared by 2 overlapping peptides, was the reactive region for clones from two DRB1\*13 patients, one carried DRB1\*1301 and one DRB1\*1302.
- Two distinct epitopes were defined in the peptide region spanning 251 to 270, and this 20-mer bound with very high affinity to DRB1\*1302 DRB1\*1301 and DRB1\*1302 would be expected to have very similar binding properties.

**HXB2 Location** p24 (129–143)

Author Location Gag (261–275 HXB-2)

Epitope IYKRWIILGLNKIVR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** DRB1\*1303, DRB1\*1502, DRB3\*0101,

DRB5\*0102, DQB1\*0301, DQB1\*0601

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ 

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 2/22 patients responded to this peptide.

**HXB2 Location** p24 (129–148) **Author Location** p24 (261–280 IIIB)

Epitope IYKRWIILGLNKIVRMYSPT

Epitope name P27

**Immunogen** in vitro stimulation or selection

Species (MHC) human

Donor MHC DR4, DR7, DQ2, DQ3

**Keywords** immunodominance, Th1, Th2, TCR usage **References** Venturini *et al.* 2002

- PBMC from a seronegative donor, the healthy brother of a pair
  of monozygotic twins discordant for HIV-1 infection, were
  used to generate HIV-1 Gag-specific CD4+ T-cell clones by
  in vitro immunization with HIV-1 overlapping 20mer peptides
  spanning p55. Six clones were generated by limiting dilution.
  All reacted with p24 except one which recognized a p24 peptide
  and a p6 peptide. All CD4+ T cell clones were HLA class II
  DR restricted.
- Clone 74 recognized two peptides including this one with a Th1 response using TCR V $\beta$  13 (13s1); it required 200 ng/ml (100 nM) and 1 µg/ml (0.5 µM) for stimulation by peptides 480-500 and 261-280, respectively. Sequencing TCR V $\beta$  regions of colonies from clone 74 suggested this was a clonal population.

**HXB2 Location** p24 (131–145)

**Author Location** Gag (298–312)

Epitope KRWIILGLNKIVRMY

Epitope name Gag 298

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

**Keywords** subtype comparisons

References Wilson et al. 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.
- This epitope binds thirteen HLA-DR alleles: DRB4\*0101, DRB5\*0101, DRB1\*0901, DRB1\*0802, DRB1\*0701, DRB1\*1302, DRB1\*1201, DRB1\*1101, DRB1\*0405, DRB1\*0401, DRB\*0301, DRB1\*1501 and DRB1\*0101, with an IC $_{50}$  threshold below 1,000 nM.
- This epitope sequence is conserved in 94% of clade B isolate.
- 8/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 non-responder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

**HXB2 Location** p24 (131–145)

**Author Location** p24 (298–312)

Epitope KRWIILGLNKIVRMY

Epitope name Gag3

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Country United Kingdom.

Assay type proliferation, Intracellular cytokine staining

**Keywords** supertype, rate of progression

References Boaz et al. 2003

Proliferative and cytokine (IFNgamma and II-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+

naive.

• Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs. Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.

**HXB2 Location** p24 (131–145) Author Location p24 (265–279 SF2)

Epitope KRWIILGLNKIVRMY

Immunogen vaccine

*Vector/Type:* virus-like particle (VLP) Strain: B clade SF2 HIV component: p24

Species (MHC) macaque

References Mills et al. 1990

• Responses to 3 T-cell and multiple linear B-cell epitopes were found in vaccinated macaques – epitope response defined by T-cell clone.

HXB2 Location p24 (131–152) Author Location p24 (263–284 SF2)

Epitope KRWIILGLNKIVRMYSPTSILD

Immunogen HIV-1 infection

Species (MHC) human

References Rosenberg et al. 1997

- Low viral load correlated with strong HIV-1-specific proliferative response.
- A proliferative response to this epitope was detected in two long term survivors.

**HXB2 Location** p24 (133–143) Author Location Gag (265–275)

Epitope WIILGLNKIVR

Epitope name Gag 14.2

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT, Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining Keywords subtype comparisons, variant crossrecognition or cross-neutralization, memory

cells

References Amara et al. 2005

- Clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better-conserved for CD8 than CD4 T cells.
- 3/5 CD8 epitopes and 2/8 CD4 epitopes were conserved across multiple HIV-1 clades. All 5 CD8, and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation.
- The similar reported human epitope in this case is KRWIIL-GLNKIVRMY, which is presented by 13 HLA-DR alleles.

T cell declined to <500 after 15 years. Patients were treatment• The response elicited to the B clade epitope WIILGLNKIVR cross-reacts with the CRF02 AG form WIvLGLNKIVR, WI-ILGLNKIVR is mostly conserved across other clades.

HXB2 Location p24 (133–144)

**Author Location** p24 (133–144 B Consensus)

Epitope WIILGLNKIVRM

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101, DRB1\*1101. DRB1\*1302, DRB1\*1501, DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This is the core of the peptide and was very often recognized, with responses by 25% of the study group. The core epitope, WIILGLNKIVRM, could bind 5/8 HLA-DR molecules tested.

**HXB2 Location** p24 (133–147)

Author Location Gag (265–279 HXB-2)

Epitope WIILGLNKIVRMYSP

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC DRB1\*1302, DRB1\*1501, DRB3\*0301,

DRB5\*0101, DQB1\*0602, DQB1\*0604; DRB1\*1303, DRB1\*1502, DRB3\*0101, DRB5\*0102, DQB1\*0301, DQB1\*0601

Country United States.

Assay type CD4 T-cell Elispot - IFNγ

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 2/22 patients responded to this peptide.

HXB2 Location p24 (133-147)

**Author Location** 

**Epitope WIILGLNKIVRMYSP** 

Epitope name G067

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine assav

Keywords memory cells References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-y, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

**HXB2 Location** p24 (133–150)

**Author Location** p24 (133–150 B Consensus)

**Epitope** WIILGLNKIVRMYSPTSI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101, DRB1\*0401,

> DRB1\*0405, DRB1\*0701, DRB1\*1101, DRB1\*1302, DRB1\*1501, DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFNy, Intracellular cytokine staining, HLA binding

Keywords supervised treatment interruptions (STI), rate of progression, immunodominance, • 8 of 24 HIV+ individuals responded to this epitope. acute/early infection

#### References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 25% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.
- The peptides that were recognized by the most people were able to bind broadly to multiple HLA-DR molecules. This peptide showed very high cross-reactive binding capacity, and bound to 8/8 common HLA-DR molecules.

**HXB2 Location** p24 (135–145)

**Author Location** p24 (135–145 B Consensus)

**Epitope ILGLNKIVRMY** 

Subtype B

Immunogen HIV-1 infection

(DRB1\*0401, DRB1\*1302, Species (MHC) human DRB1\*1501)

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cytokine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This is the core of the peptide and was very often recognized, with responses by 25% of the study group. The core epitope, ILGLNKIVRMY, could bind 3/8 HLA-DR molecules tested.

HXB2 Location p24 (135-154)

**Author Location** p24 (267–286)

Epitope ILGLNKIVRMYSPTSILDIR

Immunogen HIV-1 infection

Species (MHC) human

References Adams et al. 1997

- · One of four immunogenic Gag peptides used in study of the proliferative response to p24.
- Improved assay system (increase in culture time to 8 days and addition of IL-2 to cultures) gave increased detection of proliferative response.

**HXB2 Location** p24 (137–151)

Author Location Gag (269–285 HXB-2)

Epitope GLNKIVRMYSPTSIL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC DRB1\*1303, DRB1\*1502, DRB3\*0101, DRB5\*0102, DQB1\*0301, DQB1\*0601

Country United States.

Assay type CD4 T-cell Elispot - IFNγ

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 1/22 patients responded to this peptide.

HXB2 Location p24 (139-148)

**Author Location** p24 (271–280 HZ321)

**Epitope** NKIVRMYSPT

Subtype AG

Immunogen vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)) Strain: AG recombinant HZ321 HIV component: virus Adjuvant: Incomplete Freund's Adjuvant (IFA), CpG immunostimulatory sequence (ISS)

Species (MHC) macaque

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cytokine staining

Keywords vaccine-induced epitopes, adjuvant comparison, vaccine antigen design

References Silvera et al. 2004

• Macaques were immunized with gp120-depleted HIV-1 together with incomplete Freund's adjuvant and CpG-ODN. All four immunized animals had high anti-p24 antibody titers, while three animals showed HIV-1-specific CD4+ and CD8+ T-cell responses. This is one of two CD4+ T-cell epitopes in Gag that was mapped.

**HXB2 Location** p24 (139–157) Author Location p24 (271–290 IIIB) Epitope NKIVRMYSPTSILDIRQGP

Epitope name P28

Immunogen in vitro stimulation or selection

Species (MHC) human (DR4)

Donor MHC DR4, DR7, DQ2, DQ3

Keywords immunodominance, Th1, Th2, TCR usage

References Venturini et al. 2002

- PBMC from a seronegative donor, the healthy brother of a pair of monozygotic twins discordant for HIV-1 infection, were used to generate HIV-1 Gag-specific CD4+ T-cell clones by in vitro immunization with HIV-1 overlapping 20mer peptides spanning p55. Six clones were generated by limiting dilution. All reacted with p24 except one which recognized a p24 peptide and a p6 peptide. All CD4+ T cell clones were HLA class II DR restricted.
- Clone 6 recognized three peptides including this one with a Th1 response using TCR V $\beta$  6 (6s5A1N1). Sequencing TCR V $\beta$ regions of colonies from clone 6 suggested this was a clonal population. Assays using different peptide concentrations suggest that this peptide, 271-290, contains the main epitope of this clone. Upon activation, clone 6 was observed to induce a cytopathic effect in the adherent layer of fibroblasts expressing HLA DR4W14 and -W15. Clone 6 was activated in response to vaccinia virus Gag-infected B-LCL, so it could recognize naturally processed epitopes.
- Clone 37 recognized this peptide sequence with a Th2 response using TCR V $\beta$  3, and was a homogeneous T-cell population. This clone was only activated by peptide, not by processed protein from vaccinia virus Gag-infected B-LCL.
- Clone 97 recognized this peptide sequence with a using TCR  $V\beta$  9 and 14; the two TCR receptors used indicates this limiting dilution represents a mixed population. This clone was only activated by peptide, not by processed protein from vaccinia virus Gag-infected B-LCL.

**HXB2 Location** p24 (140–148) **Author Location** p24 (272–280 HXB2) Epitope KIVRMYSPT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101)

Donor MHC DRB1\*0101, DRB1\*1501, DO5, DO1, DR51 Assay type proliferation, T-cell Elispot, Intracellular cytokine staining

Keywords HAART, ART

References Boritz et al. 2003

- · HIV infected individuals with advanced disease often have only weak or undetectable HIV specific Th responses, and HAART can sometimes restore Th HIV recognition in such cases. The repertoire of Th responses to p24 in an individual who responded to HAART after having a CD4+ T-cell count of 0 cells/ul was determined. Eleven clonotypes were found among 13 clones, recognizing eight distinct epitopes, with a range of MHC affinities and functional avidities. Multiple Gag p24 peptides were recognized using CD4 Elispot assays based on samples from six additional HAART-treated CD4 T-cell-reconstituted subjects.
- The Th clone that recognized this epitope utilized TCR  $V\beta$ 5.2.

**HXB2 Location** p24 (141–156) **Author Location** p24 (273–287)

Epitope IVRMYSPTSILDIRQC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

- Epitope elicits a primary proliferative response in PBMC from uninfected donors.
- Matches 3/3 anchor residues for HLA DR: IVRMYSPTS

**HXB2 Location** p24 (141–158)

**Author Location** p24 (141–158 B Consensus)

Epitope IVRMYSPTSILDIRQGPK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 17% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p24 (141–158) Author Location p24 (141–158)

Epitope IVRMYSPTSILDIRQGPK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

responses.

Country Netherlands.

Assay type Cytokine production References Geels et al. 2006

- The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th
- · Autologous sequences corresponding to known and predicted Th epitopes were analyzed. IVRMYSPTSILDIRQGPK had fixation of 1 mutation (IVRMYSP[t/v]SILDIRQGPK) in 1 of the patients.

**HXB2 Location** p24 (146–160)

Author Location p24 (278–292 IIIB, B10)

Epitope SPTSILDIRQGPKEP

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

**HXB2 Location** p24 (149–168)

Author Location p24 (281–300 IIIB)

Epitope SILDIROGPKEPFRDYVDRF

Epitope name P29

Immunogen in vitro stimulation or selection

Species (MHC) human (DR4)

Donor MHC DR4, DR7, DQ2, DQ3

Keywords immunodominance, Th1, Th2, TCR usage

References Venturini et al. 2002

- PBMC from a seronegative donor, the healthy brother of a pair of monozygotic twins discordant for HIV-1 infection, were used to generate HIV-1 Gag-specific CD4+ T-cell clones by in vitro immunization with HIV-1 overlapping 20mer peptides spanning p55. Six clones were generated by limiting dilution. All reacted with p24 except one which recognized a p24 peptide and a p6 peptide. All CD4+ T cell clones were HLA class II DR restricted.
- Clone 6 recognized three peptides including this one with a Th1 response using TCR V $\beta$  6 (6s5A1N1). Sequencing TCR V $\beta$ regions of colonies from clone 6 suggested this was a clonal population. Assays using different peptide concentrations suggest that this peptide does not carry the main epitope of this clone.

**HXB2 Location** p24 (150–169)

Author Location p24 (282–301)

Epitope ILDIRQGPKEPFRDYVDRFY

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

**HXB2 Location** p24 (151–166)

**Author Location** p24 (283–297)

Epitope LDIRQGPKEPFRDYVC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

• Epitope elicits a primary proliferative response in PBMC from uninfected donors.

HXB2 Location p24 (153-167)

**Author Location** 

Epitope IRQGPKEPFRDYVDR

Epitope name G072

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine assay

Keywords memory cells

References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-γ, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

**HXB2 Location** p24 (155–177)

**Author Location** p24 (287–309)

Epitope QGPKEPFRDYVDRFYKTLRAEQA

Immunogen vaccine

Vector/Type: peptide

Species (MHC) mouse

References Nakamura et al. 1997

- Mice immunized with this peptide generated proliferative responses, CTLs and antibodies.
- This immunogenic domain is from a highly conserved region of p24.

HXB2 Location p24 (156-170)

Author Location p24 (288–302 IIIB, B10)

Epitope GPKEPFRDYVDRFYK

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

**HXB2 Location** p24 (156–173)

**Author Location** p24 (156–173 B Consensus)

Epitope GPKEPFRDYVDRFYKTLR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assav type CD4 T-cell Elispot - IFNγ, Intracellular cytokine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNy EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location p24 (156-174)

Author Location p24 (287–306)

Epitope QPKEPFRDYVDRFYKTLRA

Immunogen HIV-1 infection

Species (MHC) human

References Adams et al. 1997

- One of four immunogenic Gag peptides used in study of the proliferative response to p24.
- Improved assay system (increase in culture time to 8 days and addition of IL-2 to cultures) gave increased detection of proliferative response.

**HXB2 Location** p24 (157–165)

Author Location p24 (289–297 HXB2)

Epitope PKEPFRDYV

Subtype B

Immunogen HIV-1 infection Species (MHC) human (DQ5)

**Donor MHC** DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51

Assay type proliferation, T-cell Elispot, Intracellular cy-

tokine staining

Keywords HAART, ART

References Boritz et al. 2003

· HIV infected individuals with advanced disease often have only weak or undetectable HIV specific Th responses, and HAART can sometimes restore Th HIV recognition in such cases. The repertoire of Th responses to p24 in an individual who responded to HAART after having a CD4+ T-cell count of 0 cells ul was determined. Eleven clonotypes were found among 13 clones, recognizing eight distinct epitopes, with a range of MHC affinities and functional avidities. Multiple Gag p24 peptides were recognized using CD4 Elispot assays based on samples from six additional HAART-treated CD4 T-cell-reconstituted subjects.

**HXB2 Location** p24 (161–180)

Author Location Gag (293–312 SF2)

Epitope FRDYVDRFYKTLRAEQASQD

Immunogen vaccine

*Vector/Type:* Listeria monocytogenes Strain: B clade SF2 HIV component: p24

Species (MHC) mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

Keywords Th1

References Mata & Paterson 1999

- Listeria monocytogenes is an intracellular bacterium that lives in the cytoplasm and generates a cell-mediated immune re-
- L. monocytogenes vaccine expressing HIV-1 p24 protein (Lm-Gag) was used to stimulate gag specific CD4+ T cell proliferative responses in BALB/c(H-2d) and C57BL/6(H-2b) mice.
- 2/3 reactive p24 peptides (out of 22 overlapping peptides that span p24) were recognized by both murine strains; this peptide stimulated a response in both BALB/c and C57BL/6 mice.
- The proliferative response is due to CD4+ IFNγ-producing cells, a Th1 response.

**HXB2 Location** p24 (161–180)

Author Location p24 (293–312 HXB2)

Epitope FRDYVDRFYKTLRAEQASQD

Subtype B

Immunogen vaccine

Vector/Type: Listeria monocytogenes Strain: B clade HXB2 HIV component:

Gag

Species (MHC) mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

References Mata & Paterson 1999

- T-cells from 5 of 21 HIV+ individuals responded to this epitope. BALB/c and C57BL/6 mice were immunized with Listeria monocytogenes (Lm-Gag) expressing HIV-1 HXB2 Gag.
  - L. monocytogenes is a gram-positive bacteria that enters the macrophage on phagocytosis and lives in the cytoplasm; secreted L. monocytogenes antigens are processed and presented by both class I and class II pathways.
  - The class II Th response was probed using 20mer peptides that overlapped by 10; the peptides VHQAISPRTL-NAWVKVVEEK and FRDYVDRFYKTLRAEQASQD were recognized in H-2b and H-2d mice.

**HXB2 Location** p24 (163–175)

Author Location Gag (295–307)

Epitope DYVDRFYKTLRAE

Immunogen HIV-1 infection

Species (MHC) human (DR0101)

Assay type Cytokine production, proliferation, Tetramer

binding, CD4 T-cell Elispot - IFNγ

Keywords HAART, ART, supervised treatment interruptions (STI)

References Iyasere et al. 2003

• Fifteen patients recieving HAART with strong CD4+ proliferative responses to HIV antigens while on therapy were examined, to see the effects of viremia on these responses during treatment interruptions. Increased viremia occurred in 12/15 patients during at least one treatment interruption. Anti-HIV proliferative responses were inhibited during viremia, but IFNgamma production to Gag, Pol, and Nef peptide pools were maintained.

IL-2 production diminished during viremia, and exogenous IL-2 revived *in vitro* proliferation of HIV-specific T-cells to Gag or Pol DR0101 epitopes in a tetramer, as well as Gag-specific total CD4 T-cell responses.

HXB2 Location p24 (163–177)

Author Location p24 (295–309)

Epitope DYVDRFYKTLRAEQA

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*1302)

Keywords HAART, ART

References Blankson & Siliciano 2001; Malhotra *et al.* 2001

- The DRB1\*13-DQB1\*06 haplotype is associated with maintained viral suppression after HAART 7/7 early-treated DRB1\*13-DQB1\*06 positive people, but only 3/14 (21%) of those who did not have DRB1\*13-DQB1\*06, maintained viral suppression for 18 months.
- PBMC from individuals with the haplotype DRB1\*13-DQB1\*06 displayed increased IFNγ secretion and stronger proliferative responses against p24 80 weeks post treatment.
- DRB1\*13-DQB1\*06 was also found to be enriched among long-term non-progressors (it was in 9/18 versus, versus 21% of the general population)
- This epitope was mapped with truncated peptides using the Elispot assay, and is highly conserved.

HXB2 Location p24 (163–177)

Author Location p24 (295–309)
Epitope DYVDRFYKTLRAEQA
Immunogen HIV-1 infection

Species (MHC) human (DRB1\*1302)
Keywords HAART, ART
References Blankson & Siliciano 2001; Malhotra et al.

• The DRB1\*13-DQB1\*06 haplotype is associated with maintained viral suppression after HAART – 7/7 early-treated DRB1\*13-DQB1\*06 positive people, but only 3/14 (21%) of those who did not have DRB1\*13-DQB1\*06, maintained viral suppression for 18 months.

- PBMC from individuals with the haplotype DRB1\*13-DQB1\*06 displayed increased IFNγ secretion and stronger proliferative responses against p24 80 weeks post treatment.
- DRB1\*13-DQB1\*06 was also found to be enriched among long-term non-progressors (it was in 9/18 versus, versus 21% of the general population)
- This epitope was mapped with truncated peptides using the Elispot assay, and is highly conserved.

HXB2 Location p24 (164–181)

Author Location p24 (164–181 B Consensus)

Epitope YVDRFYKTLRAEQASQEV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101, DRB1\*0401, DRB1\*0405, DRB1\*0701, DRB1\*1101, DRB1\*1302, DRB1\*1501, DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection

#### References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The peptides that were recognized by the most people were able to bind broadly to multiple HLA-DR molecules. This peptide showed very high cross-reactive binding capacity, and bound to 8/8 tested common HLA-DR molecules.
- This peptide was the most often recognized, with a responses by 58% of the study group.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

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HXB2 Location p24 (164–181)

Author Location p24 (164–181)

Epitope YVDRFYKTLRAEQASQEV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production
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References Geels et al. 2006

- The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.
- Autologous sequences corresponding to known and predicted Th epitopes were analyzed. YVDRFYKTLRAEQASQEV had fixation of 1 mutation (YVDRFYKTLRAEQA[s/t]QEV) in 1 of the patients.

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HXB2 Location p24 (165–179)

Author Location Gag (297–311 HXB-2)
Epitope VDRFYKTLRAEQASQ
Subtype B
Immunogen HIV-1 infection

Species (MHC) human
Donor MHC DRB1*1302, DRB1*1501, DRB3*0301, DRB5*0101, DQB1*0602, DQB1*0604; DRB1*1303, DRB1*1502, DRB3*0101, DRB5*0102, DQB1*0301, DQB1*0601; DRB1*0401, DRB1*01101, DRB3*0202, DRB4*0103, DQB1*0301; DRB1*0701,
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DOB1\*0202, DOB1\*0602

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ 

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 4/22 patients responded to this peptide.

**HXB2 Location** p24 (167–178)

Author Location p24 (167–178 B Consensus)

**Epitope** RFYKTLRAEQAS

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101, DRB1\*0401, DRB1\*0405, DRB1\*0701, DRB1\*1101,

DRB1\*1501, DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, HLA binding

Keywords supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This is the core of the peptide and was the most often recognized, with responses by 58% of the study group. The core epitope, RFYKTLRAEQAS, could bind 7/8 HLA-DR molecules tested.

**HXB2 Location** p24 (168–179)

**Author Location** p24 (168–179 B Consensus)

Epitope FYKTLRAEQASQ

Subtype B

Immunogen HIV-1 infection

DRB1\*0401, Species (MHC) human (DRB1\*0101,

DRB1\*1101, DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- DRB1\*1501, DRB4\*0103, DRB5\*0101, CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
  - Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
  - This is the core of the peptide and was the most often recognized, with responses by 58% of the study group. The core epitope, FYKTLRAEQASQ, could bind 4/8 HLA-DR molecules tested.

**HXB2 Location** p24 (168–180)

Author Location p24 (168–180 B Consensus)

**Epitope** FYKTLRAEQASQE

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101, DRB1\*0401,

DRB1\*0405, DRB1\*1101, DRB1\*1501,

DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This is the core of the peptide and was the most often recognized, with responses by 58% of the study group. The core epitope, FYKTLRAEQASQE, could bind 6/8 HLA-DR molecules tested.

**HXB2 Location** p24 (169–177)

**Author Location** p24 (169–177 B Consensus)

Epitope YKTLRAEQA

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

• CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. nitude=990) were detected in 30/36 patients.

- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- This is the core of the peptide and was most often recognized, with responses by 58% of the study group. This core epitope could bind only 1/8 of HLA-DR molecules tested.

**HXB2 Location** p24 (169–178)

**Author Location** p24 (301–310 HZ321)

**Epitope** YKTLRAEQAS

Subtype AG

Immunogen vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)) Strain: AG recombinant HZ321 HIV component: virus Adjuvant: Incomplete Freund's Adjuvant (IFA), CpG immunostimulatory sequence (ISS)

Species (MHC) macaque

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining

Keywords vaccine-induced epitopes, adjuvant compari-

son, vaccine antigen design

References Silvera et al. 2004

• Macaques were immunized with gp120-depleted HIV-1 together with incomplete Freund's adjuvant and CpG-ODN. All four immunized animals had high anti-p24 antibody titers, while three animals showed HIV-1-specific CD4+ and CD8+ T-cell responses.

**HXB2 Location** p24 (169–179) Author Location Gag (301-311) Epitope YKTLRAEQASQ Epitope name Gag 15.5

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade

Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining **Keywords** subtype comparisons, variant recognition or cross-neutralization, memory cells

References Amara et al. 2005

- Clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better-conserved for CD8 than CD4 T cells.
- 3/5 CD8 epitopes and 2/8 CD4 epitopes were conserved across multiple HIV-1 clades. All 5 CD8, and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation.
- The similar reported human epitopes in this case is DYV-DRFYKTLRAEQA. HLA restriction: DRB1\*1302.

Virus specific CD4+ T-cell responses (median=7, range of mag-OASO cross-reacts with the CRF02 AG form fKTLRAEOAtO. Other clades mostly have same substitutions in these positions ([y/f]KTLRAEQA[s/t]Q).

HXB2 Location p24 (169–183)

**Author Location** 

Epitope YKTLRAEQASQEVKN

Epitope name G076

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine

Keywords memory cells

References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN- $\gamma$ , but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

**HXB2 Location** p24 (173–187)

**Author Location** Gag (301–315 HXB-2)

Epitope RAEQASQEVKNWMTE

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Donor MHC DRB1\*1303, DRB1\*1502, DRB3\*0101, DRB5\*0102, DQB1\*0301, DQB1\*0601

Country United States.

Assav type CD4 T-cell Elispot - IFNγ

References Koeppe et al. 2006

- HIV component: Env, Gag, Protease, Rev, RT, The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
  - 1/22 patients responded to this peptide.

**HXB2 Location** p24 (175–199)

**Author Location** p17 (307–331 PV22)

**Epitope** EQASQEVKNWMTETLLVQNANPDCK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*03)

Donor MHC A29, A30, B8, B35, DRB1\*03, DRB1\*13 Keywords HAART, ART, Th1, Th2, TCR usage

References Lotti et al. 2002

- 10/49 chronically HIV-1 infected patients had low p55-Gagspecific CD4+ T cell responses prior to therapy, and these responses remained unchanged 3 and 6 months after initiation of HAART. There was no difference in level of response in those with or without a detectable p55 response.
- For one individual, patient F45 CDC stage A2, CD4+ p55 responding clones were generated. Her response was consistently strong and heterogeneous in terms of HLA restriction and Vβ usage, and some clones had a Th1 cytokine secretion profile (high IFNgamma production) while some had a Th2 profile (high IL-4 and IL-5 production). 5/10 CD4+ clones could also induce cytotoxicity.
- 4/10 clones from patient F45 had their epitopes mapped using overlapping peptides. Clone 26 recognized this peptide sequence restricted by DRB1\*03. This clone had a SI of 4.1 to p55, 5.3 to peptide, secreted high levels of IFNγ, indicative of a Th1 response, but also IL-4 and IL-5. Clone 26 had no cytotoxic activity.

**HXB2 Location** p24 (181–198) **Author Location** p24 (313–327)

Epitope VKNWMTETLLVQNANC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Bedford et al. 1997

- Epitope elicits a primary proliferative response in PBMC from uninfected donors.
- Matches 3/3 anchor residues for HLA DR: VKNWMTETL

**HXB2 Location** p24 (185–202)

Author Location p24 (185–202 B Consensus)

Epitope MTETLLVQNANPDCKTIL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFN $\gamma$  EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location p24 (201–215)

Author Location Gag (333–347 HXB-2)

**Epitope ILKALGPAATLEEMM** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ 

**References** Koeppe *et al.* 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 1/22 patients responded to this peptide.

HXB2 Location p24 (205-219)

**Author Location** Gag (337–351 HXB-2)

Epitope LGPAATLEEMMTACQ

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ

References Koeppe et al. 2006

- The study measured CD4+ T-cell responses against epitopes in Gag p17 and p24 and concurrent endogenous plasma HIV-1 RNA epitope sequence variation. No intrapatient protein sequence variation was found in identified epitopes, indicating that escape from CD4-positive T-cell responses is not a common process in vivo.
- 1/22 patients responded to this peptide.

HXB2 Location p24

Author Location p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A01, A32, B\*1410, B15; A\*3101, A68,

B\*4403, B51

Country Spain.

Assay type proliferation, CD4 T-cell Elispot - IFN $\gamma$ 

Keywords HAART, ART, supervised treatment interrup-

tions (STI)

References Arnedo-Valero et al. 2004

- T cell immune responses following STI were monitored in two chronically HIV-1 infected partners (A and B) who had contracted HIV-1 during 1992. STI induced strong transitory CD4+ and CD8+ T cell responses in both patients. The viruses remained very closely related over 10 years, despite the two individuals having different HLA types; the authors suggest the maintained similarity does not support a strong role for HLA driven HIV diversity as has been claimed in Moore et al. (Science 2002).
- During the second treatment stop, patient A developed a strong proliferative response to p24, and multiple strong CD8+ T cell repsonses to Env, Pol, Gag and Nef. This patient was able to control viral load for two years follow up without therapy. Patient B developed a very weak CD4+ T cell response against

p24 during breaks in therapy, and had CD8+ responses to two epitopes. Patient A: A01, A32, B\*1410, B15; Patient B: • HIV infected individuals with advanced disease often have A\*3101, A68, B\*4403, B51.

HXB2 Location p24 Author Location p24

**Epitope** 

Immunogen vaccine

HIV component: p24 Gag Adjuvant: Keyhole Limpit Haemocyanin (KLH)

Species (MHC) human

Country United States.

Assay type proliferation, Th support of CTL response, Delayed-type hypersensitivity (DTH)

Keywords HAART, ART, immune dysfunction

References Lange et al. 2004

- ART treated HIV-1 infected patients with strong lymphoproliferative responses to HIV p24 did not have enhanced immune responses relative to those that had low level proliferative responses. Immune function was measured by DTH to diphteria/tetanus-toxoid and Keyhole limpet hemocyanin, maturation and frequency of CD8+ T cells, frequency of CD4 and CD8+ T cells, and cytotoxic molecules on HIV specific T cells.
- A higher level of persistant viral replication in circulating CD4+ cells was associated with patients who showed high lymphoproliferative responses to HIV p24.

HXB2 Location p24 Author Location p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type proliferation, Intracellular cytokine staining

Keywords HAART, ART, immune dysfunction

References Palmer et al. 2004

• The cytokine and maturation profiles as well as the proliferative capacity of HIV-1 Gag-specific CD4+ T cells was analyzed in 4 groups of HIV-1 infected patients: HAART treated, HAART suppressed, treatment naive and untreated, slowly progressing. Measurements of Gag-specific CD4+ T cell maturation, proliferation and plasma viremia indicate that virologic control is impaired due to HIV-1 affects on the maturation profiles of CD4+ T cells.

## III-B-3 Gag p2p7p1p6 Helper/CD4+ **T-cell epitopes**

**HXB2 Location** p2p7p1p6 (18–37)

Author Location p24 (384–400 HXB2)

Epitope GNFRNQRKIVKCFNCGKEGH

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DR15/51)

**Donor MHC** DRB1\*0101, DRB1\*1501, DQ5, DQ1, DR51

Assay type proliferation, T-cell Elispot, Intracellular cy-

tokine staining

Keywords HAART, ART

References Boritz et al. 2003

- only weak or undetectable HIV specific Th responses, and HAART can sometimes restore Th HIV recognition in such cases. The repertoire of Th responses to p24 in an individual who responded to HAART after having a CD4+ T-cell count of 0 cells/ul was determined. Eleven clonotypes were found among 13 clones, recognizing eight distinct epitopes, with a range of MHC affinities and functional avidities. Multiple Gag p24 peptides were recognized using CD4 Elispot assays based on samples from six additional HAART-treated CD4 T-cell-reconstituted subjects.
- The two Th clones that recognized this epitope utilized TCR  $V\beta 2$  and  $B\beta 8.1$ .

**HXB2 Location** p2p7p1p6 (22–32)

Author Location Gag (385-395)

Epitope NQRKIVKCFNC

Epitope name Gag 20.2 Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT, Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining

Keywords subtype comparisons, memory cells

References Amara et al. 2005

- Clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better-conserved for CD8 than CD4 T cells.
- 3/5 CD8 epitopes and 2/8 CD4 epitopes were conserved across multiple HIV-1 clades. All 5 CD8, and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation.
- The similar reported human epitope in this case is GNFRN-QRKIVKCFNCGKEGH. HLA restriction: DR15/51.
- The response elicited to the B clade epitope NORKIVKCFNC does not cross-react with the CRF02\_AG form gQR-IIKCFNC. The epitope is highly variable across other clades.

**HXB2 Location** p2p7p1p6 (30–44)

**Author Location** p15 (393–407 IIIB, B10)

**Epitope** FNCGKEGHTARNCRA

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

**HXB2 Location** p2p7p1p6 (37–52)

**Author Location** p15 (37–52 B Consensus)

**Epitope** HIAKNCRAPRKKGCWK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNy EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 14% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p2p7p1p6 (55–69)

Author Location p15 (418–432 IIIB, B10)

Epitope KEGHQMKDCTERQAN

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

 $\textbf{HXB2 Location} \ p2p7p1p6 \ (60-74)$ 

**Author Location** p15 (423–437 IIIB, B10)

Epitope MKDCTERQANFLGKI

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

**HXB2 Location** p2p7p1p6 (66–81)

**Author Location** p15 (66–81 B consensus)

Epitope RQANFLGKIWPSHKGR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101, DR01\*0401, DRB1\*0405, DRB1\*1101, DRB1\*1302,

DRB1\*1501, DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- Assay type CD4 T-cell Elispot IFNγ, Intracellular cyto• CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
  - This peptide was recognized by 28% of the study group.
  - Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
  - The peptides that were recognized by the most people were able to bind broadly to multiple HLA-DR molecules. This peptide showed high cross-reactive binding capacity and bound to 7/8 tested HLA-DR molecules.
  - The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p2p7p1p6 (72–89)

Author Location p15 (72–89 B Consensus)

Epitope GKIWPSHKGRPGNFLQSR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 17% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p2p7p1p6 (76–83)

**Author Location** p24 (439–446 LAI)

Epitope PSYKGRPG

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

- Stimulates T-cell proliferation in HIV-infected donors.
- Schrier lists this peptide as p24(439-446), but because of the numbering used for Gag epitopes, we placed it in p2p7p1p6.

HXB2 Location p2p7p1p6 (83–97) Author Location p15 (446–460 BRU) Epitope GNFLQSRPEPTAPPA

Immunogen in vitro stimulation or selection

**Species (MHC)** mouse (H-2<sup>b</sup>) **References** Vaslin *et al.* 1994

Peptide G4: could prime for in vitro immunoproliferative responses and for subsequent IgG responses.

HXB2 Location p2p7p1p6 (93–112) Author Location p15 (93–112 B Consensus) Epitope TAPPEESFRFGEETTTPSQK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Country** United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFN $\gamma$  EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 14% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p2p7p1p6 (98–112)

Author Location p15 (473–487 IIIB, B10)

**Epitope** ESFRSGVETTTPPQK **Immunogen** HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

Peptides were identified that commonly evoke T-cell responses
 50% of 90 HIV+ people had a T-cell response to this peptide.

**HXB2 Location** p2p7p1p6 (103–110) **Author Location** p24 (466–473 LAI)

**Epitope** REETTTPS

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

- Stimulates T-cell proliferation in HIV-infected donors.
- Schrier lists this peptide as p24(466-473), but it is in p2p7p1p6.

**HXB2 Location** p2p7p1p6 (106–116) **Author Location** Gag (469–479)

Epitope TTTPPQKQEPI

**Epitope name** Gag 24.3

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost Strain: B clade HIV component: Env, Gag, Protease, Rev, RT, Tat, Vpu

Species (MHC) macaque

Assay type T-cell Elispot, Intracellular cytokine staining

Keywords subtype comparisons, memory cells

References Amara et al. 2005

- Clade B DNA/MVA HIV vaccine was shown to raise a broad cross-reactive cellular immune response for peptides based on the CRF02\_AG consensus Gag in macaques. The activity was better-conserved for CD8 than CD4 T cells.
- 3/5 CD8 epitopes and 2/8 CD4 epitopes were conserved across multiple HIV-1 clades. All 5 CD8, and 4/8 of the CD4 epitopes that were recognized in vaccinated macaques have also been reported for human infections, indicating cross-species conservation. TTTPPQKQEPI was not reported for human infections.
- The response elicited to the B clade epitope TTTPPQKQEPI does not cross-react with the CRF02\_AG form ipssP-KQEPr. The epitope is highly variable across other clades.

**HXB2 Location** p2p7p1p6 (111–127)

Author Location p15 (111–127 B Consensus)

Epitope QKQEPIDKELYPLASLR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFN $\gamma$  EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 17% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** p2p7p1p6 (117–137) **Author Location** Gag p6 (480–500 IIIB)

**Epitope** DKELYPLTSLRSLFGNDPSSQ **Immunogen** in vitro stimulation or selection

Species (MHC) human

Donor MHC DR4, DR7, DQ2, DQ3

Keywords immunodominance, Th1, Th2, TCR usage

References Venturini et al. 2002

- PBMC from a seronegative donor, the healthy brother of a pair of monozygotic twins discordant for HIV-1 infection, were used to generate HIV-1 Gag-specific CD4+ T-cell clones by *in vitro* immunization with HIV-1 overlapping 20mer peptides spanning p55. Six clones were generated by limiting dilution. All reacted with p24 except one which recognized a p24 peptide and a p6 peptide. All CD4+ T cell clones were HLA class II DR restricted.
- Clone 74 recognized two peptides, including this one, with a Th1 response using TCR V $\beta$  13 (13s1); it required 200 ng/ml (100 nM) and 1 µg/ml (0.5 µM) for stimulation by peptides 480-500 and 261-280, respectively. Sequencing TCR V $\beta$  regions of colonies from clone 74 suggested this was a clonal population. Clone 74 was activated in response to vaccinia virus Gag-infected B-LCL, so it could recognize naturally processed epitopes.

**HXB2 Location** p2p7p1p6 (118–137)

**Author Location** p15 (118–137 B Consensus)

Epitope KELYPLASLRSLFGNDPSSQ

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFN $\gamma$  EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 17% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

### III-B-4 Gag Helper/CD4 + T-cell epitopes

**HXB2 Location** Gag

Author Location p24 (IIIB)

**Epitope** 

Immunogen in vitro stimulation or selection

Species (MHC) human (A\*0201)

Keywords dendritic cells

References Engelmayer et al. 2001

- Recombinant canarypox virus vector containing HIV-1 sequences, upon infection of mature dendritic cells, can trigger specific lysis in vitro by T-cells from HIV-1 infected individuals at levels comparable to the response seen to HIV carried in vaccinia vectors.
- Recombinant canarypox virus vector containing HIV-1 sequences can also stimulate HIV-specific IFNγ CD4+ helper T cell responses to Gag from bulk or purified CD4+ T cells.

**HXB2 Location** Gag

Author Location p55

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*13, DRB1\*03)

**Donor MHC** A29, A30, B8, B35, DRB1\*03, DRB1\*13

**Keywords** HAART, ART, Th1, Th2, TCR usage

References Lotti et al. 2002

- 10/49 chronically HIV-1 infected patients had low p55-Gagspecific CD4+ T cell responses prior to therapy, and these responses remained unchanged 3 and 6 months after initiation of HAART. There was no difference in level of response in those with or without a detectable p55 response.
- For one individual, patient F45 CDC stage A2, CD4+ p55 responding clones were generated. Her response was consistently strong and heterogeneous in terms of HLA restriction and  $V\beta$  usage. Two clones were DRB1\*13 restricted and used TCR  $V\beta$  17+19 or 5.1. Three clones were DRB1\*03 restricted and used TCR  $V\beta$  22. Some clones had a Th1 cytokine secretion profile (high IFNgamma production) while some had a Th2 profile (high IL-4 and IL-5 production).

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Qiu et al. 2000

- Mice were injected with plasmid DNA at 0, 2 and 4 weeks and lymphocyte proliferation was measured after 6 weeks with recombinant p24 protein.
- Secreted HIV-1 Gag expression vectors generated a stronger response than standard Gag or cytoplasmic Gag expression vectors
- IFN
   γ levels were increased compared to an undetectable IL-4 response.
- CTL levels were also increased in secreted Gag expression vaccination studies.

**HXB2 Location** Gag **Author Location** Gag

**Epitope** 

Immunogen vaccine

Vector/Type: DNA, DNA with protein boost Strain: B clade LAI HIV component: Gag,

Nef, Tat Adjuvant: IL-18

**Species (MHC)** mouse (H-2<sup>d</sup>) **Keywords** Th1, Th2

References Billaut-Mulot et al. 2001

- DNA vaccinated BALB/c mice primed and boosted with a multiepitopic vaccine with IL18 showed lymphoproliferative responses 7 weeks post immunization.
- Strong but non-lasting HIV-specific CTL responses were detected by a Cr-release assay and DNA prime + DNA boost was more effective than DNA prime + protein boost.
- Immunization with either the multiepitopic DNA or with the mixed DNA vaccine resulted in Th1 cytokines production (IL-2 and IFN $\gamma$ ) in spleen cell cultures stimulated by Tat and Gag, while Th2 cytokines IL-4 and IL-10 production was not detectable.
- Co-administration of IL18 increased T-cell responses but decreased anti-HIV antibody levels.

**HXB2 Location** Gag

**Author Location** p24

**Epitope** 

Immunogen vaccine

Vector/Type: coxsackievirus HIV compo-

nent: p24 Gag

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Halim et al. 2000

- An avirulent rec coxsackievirus (CB4-P) construct was generated that can express p24 Gag sequences CB4-P is attenuated even in immunodeficient mice and T help responses can be elicited from peptides embedded in a surface loop of the VP1 capsid.
- This paper describes the vaccine strategy and generation of constructs, and employs amino-terminal fusion of Gag sequences to the viral polyprotein with subsequent cleavage to elicit CTL responses via MHC class I presentation in BALB/c mice.

**HXB2** Location Gag

Author Location gp120 (V3) and p24 (IIIB, MN, BH10)

Epitope Subtype A, B Immunogen vaccine

Vector/Type: virus-like particle (VLP) Strain: A clade UG5.94UG018, B clade IIIB

HIV component: Gag, gp120

**Species (MHC)** mouse (H-2<sup>d</sup>)

**Keywords** subtype comparisons

References Buonaguro et al. 2002

- Different HIV strains were used for different regions: gp120 A clade UG5.94UG018; Gag HIV-1 IIIB
- BALB/c mice were given intraperitoneal immunization in the absence of adjuvants with virus-like particles (VLPs) expressing recombinant subtype A gp120 and Pr55gag.
- High dose-independent humoral responses were elicited against both gp120 and p24 peptides, and CTL responses were observed against target cells carrying vaccinia expressed gp120 and Gag.

**HXB2** Location Gag

Author Location Gag (HXB2)

Epitope Subtype B

Immunogen vaccine

Vector/Type: Listeria monocytogenes Strain: B clade HXB2 HIV component:

Gag

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

Keywords Th1

References Mata et al. 2001

- BALB/c and C57BL/6 mice were immunized with *Listeria monocytogenes* (Lm-Gag) expressing HIV-1 HXB2 Gag and mice were challenged with vaccinia expressing Gag.
- *L. monocytogenes* is a gram-positive bacteria that enters the macrophage on phagocytosis and lives in the cytoplasm; secreted *L. monocytogenes* antigens are processed and presented by both class I and class II pathways.
- CD4+ Th1 T-cells mediated the Gag specific immunological protection in mice immunized with Lm-Gag and challenged with vaccinia-Gag.
- Gag-specific CTL may enhance viral clearance via IFN
   γ secretion, but are not essential for immunity.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen vaccine

Vector/Type: Listeria monocytogenes HIV

component: Gag

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

Keywords review, Th1

References Mata & Paterson 2000

- BALB/c and C57BL/6 mice were immunized with Listeria monocytogenes (Lm-Gag) expressing HIV-1 HXB2 Gag and mice were challenged with vaccinia expressing Gag.
- *L. monocytogenes* is a gram-positive bacteria that enters the macrophage on phagocytosis and lives in the cytoplasm; secreted *L. monocytogenes* antigens are processed and presented by both class I and class II pathways.
- This article is a review of *L. monocytogenes* biology and its potential as a vaccine vector for HIV, comparing to other vector systems, and discussing CD4+ Th1 T-cell-mediated Gag specific immunological protection in mice and the Gag CTL response.

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: virus-like particle (VLP) HIV

component: p17 Gag, p24 Gag

Species (MHC) human

References Kelleher et al. 1998b

- Immunization of HIV+ people with a p24-VLP virus-like particle did not significantly impact CD4+ lymphocyte count, viral load, or p24 antibody titre.
- Immunization with p24-VLP showed a modest, short-lived increased proliferative response to p24.

**HXB2 Location** Gag Author Location p24 **Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)), protein Strain: AG recombinant HZ321 HIV component: gp120 depleted virus, p24 Gag

Species (MHC) human

References Maino et al. 2000

- 18 HIV-1-seropositive patients with a low frequency or no detectable CD4+ T cell response to HIV-1 antigen received an HIV-1 immunogen consisting of 10 units of native p24 and 100 ug of HZ321, a gp120 depleted antigen.
- Using flow-cytometric methods, HIV-1 specific CD4+ T cells were shown to increase in response to immunization – in many patients significant enhancement was observed after a single immunization.
- The frequency of CD4+ T cells expressing cytokines in response to antigen by FACS was correlated with a lymphoproliferation assay.

**HXB2 Location** Gag Author Location p24 **Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, supervised treatment interruptions (STI)

References Ruiz et al. 2000

- Structured treatment interruption in chronically infected patients allowed recovery of p24 Th proliferative responses after HAART therapy discontinuation in 2/12 patients.
- The Th response to p24 was identified during peak viremia in one patient, while in the second it was noted when viremia was controlled after restarting antiviral therapy.

**HXB2 Location** Gag Author Location p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Lori et al. 1999

- Ten patients with acute, pre-seroconversion HIV-1 infections were treated with didanosine, indinavir and hydroxyurea - this treatment is associated with normalization of immune parame-
- A vigorous HIV-specific Th response (stimulation index greater than 8) was observed in 7/8 patients treated before complete WB seroconversion, but in only 1/5 controls treated after sero-
- Vigorous Th responses were detected as early as 34 days after treatment begin.
- Patients treated prior to seroconversion had no loss of naive CD4 T lymphocytes, recovery of up to 35% of the naive CD8 cells in several weeks, and a reduced latent viral reservoir.

**HXB2 Location** Gag Author Location p24 **Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, supervised treatment interruptions (STI), Th1

References Haslett et al. 2000

- 11/22 adult patients on HAART showed strong CD4+ T-cell IFNγ producing Th1 responses to HIV p24.
- The magnitude of the Th1 response correlated with previous interruptions in HAART, suggesting the interruptions primed or boosted the response.
- In contrast, the magnitude of the CD8+ CTL response did not correlate with interruptions in therapy, although a greater breadth in response was associated with interruptions in HAART.

**HXB2 Location** Gag Author Location p24 **Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: virus-like particle (VLP) HIV component: p17 Gag, p24 Gag

Species (MHC) human

References Klein et al. 1997

- Immunization of HIV+ people with a HIV-1 p17/p24 Ty viruslike particle (p24-VLP) resulted in a marginal, short-lived increased proliferative response to p24 and p17 and a transient elevation in viral load.
- Two of four subjects that received 500 or 1000 ug of p24-VLP had an increase in gag-specific CTL.

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)) Strain: AG recombinant HZ321 HIV component: gp120 depleted

Species (MHC) human

**Keywords** subtype comparisons

References Moss et al. 1998

• Immunization with gp120 depleted HZ321 virus (REMUNE<sup>TM</sup>) triggered an increase in lymphocyte proliferative response to native p24, a clade B virus and clade E viral antigens – Z321 is clade A in env and clade G in gag. Moss et al. [1998]

**HXB2 Location** Gag Author Location p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Rosenberg et al. 1999

- This paper reviews the role of T-cells in viral control and HIV disease outcome.
- Strong anti-p24 lymphoproliferative responses were found in seven persons who were treated with potent anti-viral therapy during acute HIV-1 infection syndrome.

 This suggests that Th cells are part of the normal response to HIV-1 infection, but their numbers are rapidly diminished by either being infected during the peak viremia or by activationinduced cell death – if peak viremia can be controlled, a robust anti-p24 Th response can be maintained.

**HXB2 Location** Gag **Author Location** p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

**References** Rosenberg & Walker 1998

- Strong Th responses have been found in rare individuals who effectively maintain low viral loads.
- If aggressive anti-retroviral therapy is given prior to seroconversion, strong helper responses can be maintained.

HXB2 Location Gag
Author Location p17
Epitope
Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

Species (MHC) mouse

References Birk et al. 1998a

• Different p17 genes derived from the same quasispecies and expressed and purified in E. coli primed different Th 1 and Th 2 subsets in mice, depending on their H-2 type.

HXB2 Location Gag Author Location Gag Epitope

Immunogen HIV-1 infection

Species (MHC) human

References Schiller et al. 2000

- Study of parameters that might influence the performance or reproducibility of clinical Th proliferative assays.
- HIV-1 replication in vitro is unlikely to influence the assay.
- Gag proteins including p17 and possibly p7 as well as p24 perform better than p24 alone.
- Frozen samples can be used in T-proliferative assays, but with lower radiolabled thymidine incorporation.

HXB2 Location Gag Author Location Gag Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART **References** Pitcher *et al.* 1999

- In contrast to earlier studies suggesting that HIV-1 specific Th
  responses were eliminated in the early stages of infection in
  most HIV+ individuals, this paper shows using flow cytometric
  detection of antigen-induced cytokines that Th-1 CD4+ memory gag-specific Th cells are detectable in most HIV+ subjects.
- Effective anti-viral therapy reduces the frequency of these cells, presumably due to reduced antigenic stimulus.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART **References** Plana *et al.* 1998

 Patients from later stages of infection given HAART do not show restoration of HIV-1 specific Th proliferative responses.

HXB2 Location Gag Author Location Gag

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Kelleher et al. 1998a

• Env and gag Th epitopes were pooled and used to test Th proliferative responses after IL2 therapy – while IL2 therapy causes an increase in CD4+ lymphocyte count, it does not increase HIV-1 specific proliferative responses.

**HXB2 Location** Gag **Author Location** Gag (LAI)

Epitope Subtype B Immunogen vaccine

Vector/Type: DNA prime with vaccinia boost Strain: B clade LAI HIV component: Env,

Gag

Species (MHC) macaque

Keywords Th1, Th2

References Kent et al. 1998

- Priming with an HIV-DNA vaccine and boosting with a vaccinia construct induced greater levels of HIV T-cell immunity than either vaccine alone.
- The proliferative response to Env and Gag after the DNA vaccination had a mean SI of 1.5-4, but after boosting with rHIV-fowlpox virus, there was a 6-17 fold increase in the mean SI for HIV Gag and Env The Th response happened despite a fall in Ab titers, suggesting that the Th response was primarily Th1, not Th2. The CTL response was also enhanced.

**HXB2 Location** Gag

**Author Location** 

Epitope

Immunogen vaccine

*Vector/Type:* DNA, protein, virus-like particle (VLP), ISCOM

Species (MHC) macaque

Keywords Th1, Th2

References Heeney et al. 1999

- Ten different vaccine strategies were evaluated for their ability to protect from infection in a rhesus macaque model using a non-pathogenic SHIV challenge.
- Protection correlated with the magnitude of NAb responses, beta-chemokines, and a balanced Th response.
- DNA, protein+adjuvant, VLP and ISCOM vaccines were tested.

• HIV-1/ISCOMS gave the highest NAb titers, Th1 and Th2 responses, was the only vaccine formulation tested with a detectable CTL response, and gave enhanced beta-chemokine production.

**HXB2 Location** Gag

Author Location Gag/Pol (MN)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade MN HIV component: Env, Gag, Pol Adjuvant: CD80, CD86

Species (MHC) chimpanzee

References Kim et al. 1998

Co-stimulatory molecules co-expressed with an HIV-1 immunogen in a DNA vaccine used to enhance the immune response – co-expression of CD86, but not CD80, dramatically increased both HIV Env and Gag/Pol specific CTL and Th proliferative responses.

**HXB2** Location Gag

Author Location Gag/Pol (LAI, MN)

Epitope

Immunogen vaccine

Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag, gp120, gp41, Protease

Species (MHC) human

References Salmon-Ceron et al. 1999

 A live attenuated canarypox vector expressing MN gp120 and LAI gp41/gag/protease could induce CTL and a lymphoproliferative response in healthy uninfected volunteers.

**HXB2 Location** Gag

Author Location p55 (IIIB)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART **References** Zhang *et al.* 2001b

- T-helper cell proliferative responses to HIV p24, p55 and gp120 were tested in 27 patients with HIV infection vigorous responses directed at Gag were detected in ten patients, but an Env specific response was detected in only one patient.
- Untreated patients showed a negative correlation between plasma viral load and HIV p24-specific T-cell responses, and the responses could be detected after extended HAART therapy with viremia below the detection limit.

**HXB2 Location** Gag

**Author Location** p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI), kinetics, Th1

References Carcelain et al. 2001

- Repeated structured HAART therapy interruptions (STI) in 3 chronically HIV infected patients induced rapid but transient (< 3 weeks) HIV-1 specific CD4+ Th1 responses concurrently with viral rebound, as measured by proliferation assays and by IFNγ production by CD8-depleted PBMC.</li>
- Kinetics suggest that viral replication leads to rapid destruction of the HIV-specific Th1 cell response.
- HIV-specific CD8+ T-cell responses were delayed relative to the Th1 responses and were not sustained.

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Blankson et al. 2001

- 5/10 chronically HIV infected patients with low CD4+ counts who received HAART therapy and experienced immune reconstitution displayed p24, p17 and p66 T-helper CD4 proliferative responses, in contrast to 0/8 chronically HIV infected patients with high CD4+ counts at the initiation of antiretroviral treatment.
- This surprising result could be due to the low CD4 nadir patients being more likely to have thymic regeneration or a peripheral expansion of T cells.

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART

References Angel et al. 2001

- Prolonged viral suppression resulting from potent antiretroviral therapy allowed a T helper response to Gag p24 and PHA to develop in many HIV+ patients.
- At baseline, 2/41 (4.9%) subjects had a proliferative response to Gag p24, and 7/41 (17.1%) had a response to PHA, but by week 72 of therapy, 53% had a detectable response to p24 and 94% to PHA.

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Blazevic et al. 2000

 Prolonged viral suppression resulting from potent antiretroviral therapy did not allow an HIV T helper response to p24 or gp160, but Th proliferative responses to influenza, alloantigen, and PHA did develop in many HIV+ patients, and asymptomatic patients had stronger and more frequent Th response recovery than AIDS patients.

**HXB2 Location** Gag

**Author Location** Gag (SF2)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, acute/early infection

References Altfeld et al. 2001b

- Therapy provided during acute infection resulted in a narrower CTL response, stronger T help response, and a less diverse viral population than was seen in individuals treated during chronic infection.
- The breadth and specificity of the CTL response was determined using Elispot by studying 19 individuals with preseroconversion therapy (Group 1), 11 individuals with primary infection but post-seroconversion therapy (Group 2), and 10 individuals who responded to HAART given during chronic infection (Group 3), using 259 overlapping peptides spanning p17, p24, RT, gp41, gp120 and Nef.
- Individuals who were given HAART during acute or early in infection had significantly stronger proliferative responses than individuals who were chronically infected.

HXB2 Location Gag Author Location p24

Epitope

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Oxenius et al. 2000

- Patients who started therapy at acute HIV infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV-specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV-specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.
- In 3/4 responders tested p24 gave the strongest T helper response.

**HXB2 Location** Gag **Author Location** p24

**Epitope** 

Immunogen vaccine

Vector/Type: gp120 depleted whole killed virus Strain: AG recombinant HZ321 HIV component: virus Adjuvant: Complete Freund's Adjuvant (CFA), CpG immunostimulatory sequence (ISS)

Species (MHC) rat

References Moss et al. 2001

- Different HIV strains were used for different regions: subtype A env, subtype G gag
- Lewis rats simultaneously immunized with HIV-1 antigen and with immunostimulatory sequences CpG had increased Th proliferative responses, but when CpG was given as a prime prior to the injection of HIV-1 antigen it was not as effective.

HXB2 Location Gag
Author Location p24
Epitope
Immunogen vaccine

Vector/Type: gp120 depleted whole killed virus Strain: AG recombinant HZ321 HIV component: virus Adjuvant: Complete Freund's Adjuvant (CFA), CpG immunostimulatory sequence (ISS)

Species (MHC) rat

References Moss et al. 2000

- Different HIV strains were used for different regions: subtype A env, subtype G gag
- Lewis rats co-immunized with HIV-1 antigen in Freund's and with immunostimulatory sequences CpG stimulated increased IFN $\gamma$  expressing CD4+ and CD8+ T cells and anti-p24 anti-bodies relative to antigen in Freund's without CpG.

**HXB2 Location** Gag

**Author Location** p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression, Th1

References Kalams et al. 1999a

- The strength of p24 specific Gag proliferative responses (SIs) were inversely correlated with viral load in 21 ARV naive patients. The responses were Th1, IFN $\gamma$  producing. Proliferative responses against gp160 were rarely observed (only 4 cases).
- Gag specific CTL levels were correlated with Gag proliferative responses but were not correlated with viral load. 8 subjects lacked p24 specific Gag proliferative responses, and 4/8 had no CTLp to any HIV-1 antigen tested.

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART, review, rate of progression

References Kalams & Walker 1998

 This paper reviews the role of specific T cell help in many viral infections, and covers the interplay between Th, CTL and survival, and discusses briefly advantages of HAART during acute HIV infection to prevent the early decimation of the Th response in HIV infections.

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression, Th1, Th2

References Wilson et al. 2000b

- Dysfunction of HIV-1 specific proliferative responses, but not responses to other antigens, is evident in HIV-1 progressive disease.
- Vigorous HIV-1 specific responses to p24, Nef and gp120 with SI between 8-99 were seen in 6/7 long term non-progressors (LTNP), the seventh had a borderline responses. IL-2 production was seen in all cases, and IL-4 production was also evident many responses.
- None of the progressors (0/5) had HIV-1 specific proliferative responses, or IL-2 or IL-4 induction.

• Non-HIV antigens (cytomegalovirus, PPD, Staphylococcus enterotoxin B, tetanus toxoid) gave similar responses in HIV-1 + LTNP, progressors, and HIV-1 controls.

**HXB2 Location** Gag Author Location p24 **Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression, Th1 References Alatrakchi et al. 2002

- · LTNP co-infected with HCV and HIV showed higher frequencies of Th1 response to both HIV-1 p24 and HCV antigens.
- HIV-1 CD4 Th1 responses in untreated LTNP were inversely correlated with viral load.

**HXB2 Location** Gag Author Location p24 **Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART References Lange et al. 2002

- · Cross-sectional study compares CD4 T-cell count and age matched untreated HIV-1 + patients (N = 14) with patients undergoing HAART therapy (N = 14).
- The fractions of naive and memory T-cells were comparable for both groups, as were proliferative responses to non-HIV antigens. Lymphocyte proliferation responses to HIV-1 p24 were of greater magnitude in the group treated with HAART (5/10 had SI > 10, versus 1/12 in the untreated group), suggesting that ongoing viral replication impairs the anti-Gag response, and the response can be improved and restored through HAART.
- DTH responses to recall antigens were tested, and responses to C. albicans and Trichophyton were comparable in both treated and untreated patients, although patients on therapy had higher responses to mumps.

**HXB2 Location** Gag Author Location p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, subtype comparisons, escape, acute/early infection

References Fidler et al. 2002

- 37/45 patients with primary HIV infection underwent a short course of antiretroviral therapy (SCART). 29/37 patients received triple ART therapy and eight patients received four ART drugs. Initiation of SCART was effective in controlling HIV replication by ten weeks in all patients and preserving CD4+ T cell responses for up to 64 weeks after therapy.
- No induction of drug escape mutations was observed, although two individuals had escape mutations in their infecting virus at
- 34 UK infected patients were clade B infected. 11/45 subjects had non-UK acquired HIV infection, 2 were clade A, 1 was A/E, 1 was C, 1 was "untypable", the rest were B.

- Recombinant HIV-1 derived gp120, p24, p66 and overlapping peptide pools spanning Tat and Nef were employed to measure CD4 T-cell frequencies in ELISPOT assays. The strongest preservation of T helper responses 12 weeks off SCART was seen for p24-specific CD4+ T-cell responses.
- 6/8 of the untreated individuals were tested for CD4+ T-cell responses. 1 had no detectable response. 1 had detectable responses to all HIV-1 proteins tested at baseline, but this narrowed to p24 and gp120, then became undetectable by 52 weeks. 3 had detectable and persistent responses, but only to
- Post-therapy, the average spot forming cells for all proteins tested in 17/37 with 24 weeks of follow up had not declined, although the plasma viral RNA was increasing. SFU using p24 were measurable following SCART and preserved at levels comparable to baseline.

**HXB2 Location** Gag **Author Location Epitope** Subtype B

Immunogen vaccine

Vector/Type: virus-like particle (VLP) Strain: B clade IIIB HIV component: p17 Gag, p24 Gag Adjuvant: aluminum hydrox-

Species (MHC) human

Keywords rate of progression

References Klein et al. 1997; Lindenburg et al. 2002

• HIV-1 p17/p24:Ty virus-like particles therapeutic vaccination of 56 HIV-1 infected patients had no effect on disease progression, AIDS and CD4+ T-cell decline in a longitudinal study, despite some evidence suggesting it can enhance Th anti-Gag proliferative responses in HIV+ individuals Klein et al. [1997]

**HXB2 Location** Gag Author Location p24 (NY5)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection, crosspresentation by different HLA, early treatment

References Norris et al. 2001

- Gag-specific CD4+ helper T-cell clones were derived from 1 LTNP (CTS-01) and 3 individuals given therapy during acute infection, 2 before (AC-01 and AC-36) and 1 after (AC-25)
- The immunodominant response in LTNP CTS-01 was to peptide 9, and 9/10 clones derived from this patient reacted with it. Three, two, and one clones were obtained from the 3 patients given therapy. These 6 clones all reacted with different p24 peptides, and all had peptide induced proliferative responses, IFN $\gamma$  production, and cytotoxic responses. The implications of cytotoxic responses in CD4+ T-helper cells are discussed.

**HXB2 Location** Gag

Author Location p24 Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART **References** Palmer *et al.* 2002

- CD4+ T cell proliferative responses from 33 HIV-1 infected patients with HAART suppression were compared to 19 patients with active viral replication (HAART failures and HAART naive). Patients with HAART suppression showed stronger p24- and p66-specific proliferative responses compared to patient groups with active HIV-1 replication, suggesting active viral replication *in vivo* specifically reduces proliferation responses.
- No differences in the frequency of HIV-specific CD4+ T-cells that were positive for cytokine secretion in a flow cytometry assay were found in the HAART suppressed group versus the group with active viral replication.

HXB2 Location Gag

**Author Location** p24 (SF2)

**Epitope** 

Subtype B, G

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression, Th1, Th2

References Imami et al. 2002b

- 70 patients with chronic disease progression, 10 clinical nonprogressors, and 3 immunologically discordant progressors (individuals who controlled viremia but had progressive CD4+ T-cell decline) were analyzed for their T-helper cell responses to p24 and cytokine profile.
- SF2 p24 20mer peptides overlapping by 10 were used to assess the response in the different groups. At least 1/10 and up to 7/10 nonprogressors had a proliferative response with every one of the 22 p24 overlapping peptides. All peptides produced an IL-2 (Th1) response in at least one of the 10 nonprogressors. IL-4 (Th2) responses were strong, but somewhat less comprehensive as 6/22 peptides elicited no IL-4 production, and fewer IL-4 responses were see per peptide. In contrast, only 1/10 progressors had a clear proliferative and IL-2 response to 2/22 peptides, and neither one made an IL-4 response.
- The results taken together suggest that a balanced Th1/Th2 response to HIV is important for viral control in long-term non-progression.
- One immunologically discordant progressor became symptomatic while on the study. He showed a rapid decline in proliferative activity at that point, and a shift from a Th1 to a Th2 IL-4 producing response.

**HXB2 Location** Gag

**Author Location (BRU)** 

**Epitope** 

Subtype B

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade BRU HIV component: virus Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse

References Haas et al. 1991

- Of 5 mouse inbred lines tested: DBA/2 (H-2d, Ad, Ed), B10.A(4R) (H-2h4, Ak) and B10.A(5R) (H-2i5) showed particularly good CD4+ T cell proliferative responses to HIV proteins (gp160, gp120, p17, p24, Nef and RT), after vaccination with inactivated virus.
- B10.BR (H-2k, Ak, Ek) and C57BL/6 (H-2b and Ab) had weaker responses.

**HXB2 Location** Gag

Author Location Gag (III-B)

**Epitope** 

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB

HIV component: Gag

**Species (MHC)** mouse **Donor MHC** H-2<sup>d</sup>

Keywords vaccine-specific epitope characteristics, Th1

References Bojak et al. 2002a

 Codon-optimized gag gene DNA vaccines were compared to wild type by vaccination of BALB/c mice. Codon optimized DNA gave a Th1 polarized Th response, a strong antibody response that persisted from more than 20 weeks, and CTL responses, while wild-type DNA induced weak and inconsistent immune responses.

**HXB2** Location Gag

Author Location Gag (MN)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type Cytokine production, proliferation, CD4 Tcell Elispot - IFNγ, Intracellular cytokine

staining

Keywords HAART, ART, acute/early infection

References Malhotra et al. 2003

92 acute- or early-HIV infected subjects were tested for Envand Gag-specific Th responses. There was an overall low probability of detecting HIV-1-specific Th responses, Env responses were rare and (5%) and only detected very early in infection, probably reflecting a low frequency of response and impaired lymophoproliferative capacity, not viral escape. Gag-specific Th responses were observed more frequently (16%), were found throughout the acute and early infection phases, and predominated after ARV therapy.

HXB2 Location Gag

Author Location p24

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)), protein Strain: AG recombinant HZ321 Adjuvant: Incomplete

Freund's Adjuvant (IFA)

Species (MHC) human

Assay type Cytokine production, proliferation, T-cell Eli-

spot

Keywords HAART, ART, supervised treatment interrup-

tions (STI), immunotherapy

References Moss et al. 2003

• Structured treatment interruptions (STIs) were compared in individuals that had been given prior therapeutic vaccines, and those that had not. Therapeutic immunization increased gag p24 stimulated proliferative responses and MIP-1β responses prior to STIs, although total CD4 counts viral RNA levels were unchanged. Proliferative responses and chemokine induction in the vaccinated group correlated with the control of viremia during subsequent STIs.

HXB2 Location Gag Author Location Epitope

Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Assay type** proliferation, T-cell Elispot, Intracellular cytokine staining

Keywords supertype

References Papasavvas et al. 2003

- Children with full or partial viral supression along with stable CD4+ T cell counts had significantly increased levels of anti-HIV CD4+ T cell proliferative responses, and decreased CD38+ T-cells
- Preservation of high levels of CD4+ T-cells was associated with a high percentage of CD4+ naive T-cells relative to memory T-cells

HXB2 Location Gag Author Location p24 Epitope

Immunogen HIV-1 infection, vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)) Strain: AG recombinant HZ321 Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human

Assay type proliferation, T-cell Elispot, Delayed-type hypersensitivity (DTH)

Keywords HAART, ART, immunotherapy

References Robbins et al. 2003

- Augmented Th cell responses to Gag p24 were seen in five out of five chronically infected individuals who had virological control with HAART, after therapeutic immunization with REMUNE (gp120 depleted inactived virus). The magnitude of responses ranged from a 5- to 200-fold increase, with fluctuation in magnitude over time.
- There was no change in the magnitude and breadth of CTL responses, CD4 counts or percentages, or DTH responses.

**HXB2 Location** Gag **Author Location** p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Donor MHC** A1, A2, B8, B44, DR4, DR15; LTNP S24: A2, A11, B55, B57, DR4, DR13; LTNP C135:

A1, A33, B50, B57, DR7, DR13

Assay type Cytokine production, proliferation, CD8 T-

cell Elispot - IFN $\gamma$ , CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cytokine staining

**Keywords** rate of progression, immunodominance **References** Wang *et al.* 2002a

Gag Helper/CD4+ T-cell epitopes

- A 51 year old male, infected presumably in 1988, diagnosed seropositive in 1993, has remained asymptomatic and is a long term non-progressor. He had very low proviral copy number in his PBMCs with high levels of G-A hypermutation, resulting in multiple stop codons, and viral replication was not evident. He was heterozygous for the CCR5 delta 32 allele, and has undergone a variety of treatments through the years. T cell responses in this patient and in two additional LTNPs were described, and this patient had particularly intense CD4+ Th responses.
- PBMC from this patient resisted infection from CCR5, CXCR4 and dual-tropic HIV-1 strains. Purified CD4+ T cells became infected, however, without detectable cytopathic effect. CD8+ T cells were shown to protect PBMCs from infection, and this protection was not mediated by IFNgamma. Undefined CD8 T-cell secreted factors were stimulated by Gag, Pol and Nef genes introduced into target cells with vaccinia and processed through a class I pathway were responsible for the protective effect. This factor resembled CAF, the CD8+ cell antiviral factor described in Mackewicz and Levy (ARHR 8:1039, 1992)
- The CD4+ and CD8+ T-cell populations were both strongly skewed toward the CD45RO+ phenotype, many of which were terminally differentiated, CD28-, and expressed the activation markers CD38+ and HLA-DR+. Cell turnover, however, wasn't much elevated as measured by apoptosis or Ki-67+ and Bcl-2 dim expression.
- Vigorous p24-specific Th proliferative responses were observed, and 50% of CD4+ T-cells proliferated in response to p24 Gag, an extraordinary percentage. Responses were also detected against other regions in Gag, gp120 and Nef. It remains unclear how such vigorous Th responses are maintained with undetectable ongoing viral replication.
- Strong CD4+ T-cell IFNgamma Elispot responses were mapped to many peptides in Gag for this patient. T-cells from two other LTNPs were tested here, and they did not react with as many Gag peptides as the main study subject of the paper. NIH reference Gag peptide set was used, but the sequences of the reactive peptides and the precise strain was not indicated in the paper, so we could not record them in the database.
- CD8+ T cell Elispot responses to Gag, Env, Nef, and Pol were detected as well, although CTL were not prominent, consistent with undetectable viremia.
- This subject had strong NAb responses when tested using the X4 primary isolate 228 200.

HXB2 Location Gag Author Location p24 Epitope

Immunogen HIV-1 infection

Species (MHC)

Assay type proliferation Keywords HAART, ART References Sullivan *et al.* 2003

 Lymphoproliferative responses to HIV antigens p24, gp120 and Nef were enhanced in eight patients who were switched from protease inhibitors to non-nucleoside reverse transcriptase inhibitors. **HXB2 Location** Gag **Author Location** p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type Cytokine production, proliferation

**Keywords** HAART, ART **References** Hardy *et al.* 2003

 Upon initiating HAART, CD4+ T cell proliferative responses in 36 patients were restored to specific antigens, mitogens, and IL-2. Restored recall responses were largely to persistent antigens, and not to HIV-1 or new antigens, and recall responses were associated with IL-2, not IL-4 production.

HXB2 Location Gag

Author Location p24

**Epitope** 

Immunogen HIV-1 and HCV co-infection

Species (MHC) human

Assay type CD4 T-cell Elispot - IFNγ

**Keywords** HAART, ART, Th1

References Alatrakchi et al. 2004

 Treatment with IFNalpha and ribavirin induced a threefold decrease of type 1 T-helper cell frequencies specific for HIV (p24) and CMV in HIV/HCV co-infected patients undergoing HAART therapy, suggesting this therapy might negatively impact viral-specific immune responses.

**HXB2 Location** Gag

**Author Location** p24

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Country Spain.

Assay type proliferation

Keywords HAART, ART, supervised treatment interrup-

tions (STI)

References Plana et al. 2004

- Study evaluated the dynamics of CD4+ and CD8+ T-cell responses during 4 cycles of STI in 45 patients, who had early-stage, chronic HIV-1 infection. Lymphoproliferative responses (LPRs) increased between the beginning of the first STI cycle through the 4th STI, but then decreased. Viral load at the end of the 4th STI was inversely correlated with p24 LPRs, but the LPRs were transient and after 12 weeks no longer were correlated with low viral low.
- STIs can boost CTL and LPR responses, but the lack of durable T-helper responses leads to lack of long term viral control.

**HXB2** Location Gag

**Author Location** 

**Epitope** 

Subtype CRF02\_AG

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human

Country Senegal.

Assay type CD4 T-cell Elispot - IFNγ

Keywords rate of progression, variant cross-recognition

or cross-neutralization

References Zheng et al. 2004

- Gag, Env, Tat, and Nef-specific T-cell responses were evaluated in 68 HIV-1 and 55 HIV-2 infected drug naive, generally asymptomatic, infected Senegalese patients.
- HIV-1 peptides were derived from HIV-1 CRF-02 (HIV-1 A/G, AJ251056)and HIV-2 peptides spanning HIV-2 ROD (M15390).
- Gag specific responses dominated in both groups, but overall magnitude and frequencies did not correlate with viral load or CD4 counts. CD4+ Helper T-cell responses were found in only 8% of HIV-1 + people, but in 48% of HIV-2 + people, suggesting helper T cell responses may contribute to improved control of viremia in HIV-2 infected patients. Lower viral load was associated magnitude of T-cell responses in HIV-1 infection only when the T-cell responses were measured for cross-reactivity with HIV-2.

**HXB2** Location Gag

Author Location p24

**Epitope** 

Subtype A, AG, B

Immunogen HIV-1 infection

Species (MHC) human

Country Cote D'Ivoire.

Assay type Cytokine production, CD8 T-cell Elispot -

IFN $\gamma$ , CD4 T-cell Elispot - IFN $\gamma$ 

**Keywords** HIV exposed persistently seronegative (HEPS)

References Jennes et al. 2004

- Env(gp120)- and Gag(p24)-specific T helper responses were compared between HIV-exposed seronegative (ESN) and seropositive female sex workers in Africa (Abidjan, Cote d'Ivoire).
- HIV-specific CD4+ T cells were detected in both study groups; low level EliSpot responses were found in 8/40 ESN sex workers. The presence of HIV specific CD4+ T-cells was detected by flow cytometry in 3/8 (38%) in the ESN group, was associated with the frequency and not with the duration of HIV exposure. The ESN reponses were detected in women with more clients on the previous working day and more exposures per month.
- B subtype peptides were used to probe these responses because of availability, however the predominant clades circulating in the area are A and CRF02.

### III-B-5 RT Helper/CD4 + T-cell epitopes

HXB2 Location RT (36-52)

Author Location RT (36-52 BRU)

Epitope EICTEMEKEGKISKIGP

Immunogen HIV-1 infection

Species (MHC) human

References De Groot et al. 1991

 9 out of 17 humans can make strong IL2 responses to this epitope.

HXB2 Location RT (38-52)

**Author Location** RT (38–52 BRU)

**Epitope** CTEMEKEGKISKIGP

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: RT

**Species (MHC)** mouse  $(H-2^k)$ 

References De Groot et al. 1991

 T-cells from RT immunized mice have enhanced proliferative response with peptide.

HXB2 Location RT (39-53)

Author Location RT (194-208)

Epitope TEMEKEGKISKIGPE

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995a

• Protein priming induced T-cells that recognize peptide, 4 clones from a single donor recognized this peptide.

HXB2 Location RT (48–62)

**Author Location** RT (48–62 BRU)

**Epitope** SKIGPENPYNTPVFA

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: RT

**Species (MHC)** mouse (H-2<sup>k</sup>)

References De Groot et al. 1991

 T-cells from RT immunized mice have enhanced proliferative response with peptide.

HXB2 Location RT (62–77)

**Author Location** RT (62–77 BRU)

Epitope AIKKKDSTKWRKLVDF

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: RT

**Species (MHC)** mouse (H-2<sup>k</sup>)

**References** De Groot et al. 1991

 T-cells from RT immunized mice have enhanced proliferative response with peptide.

HXB2 Location RT (88-102)

**Author Location** RT (88–102 BRU)

Epitope WEVQLGIPHPAGLKK

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: RT

Species (MHC) mouse (H-2<sup>t4</sup>)

References De Groot et al. 1991

 T-cells from RT immunized mice have enhanced proliferative response with peptide.

**HXB2 Location** RT (124–138)

Author Location Pol (303–317)

Epitope FRKYTAFTIPSINNE

Epitope name Pol 303

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

**Keywords** subtype comparisons **References** Wilson *et al.* 2001

 Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.

- This epitope binds seven HLA-DR alleles: DRB1\*0901, DRB1\*0802, DRB1\*0701, DRB1\*0405, DRB1\*0401, DRB1\*1501 and DRB1\*0101, with an IC<sub>50</sub> threshold below 1,000 nM.
- This epitope sequence is conserved in 68% of clade B isolates.
- 8/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 non-responder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

**HXB2 Location** RT (124-138)

**Author Location** RT (303–317)

**Epitope** FRKYTAFTIPSINNE

Epitope name Pol1

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Country United Kingdom.

Assay type proliferation, Intracellular cytokine staining

Keywords supertype, rate of progression

References Boaz et al. 2003

- Proliferative and cytokine (IFNgamma and II-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+T cell declined to <500 after 15 years. Patients were treatment-naive.</li>
- Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs.
   Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.

HXB2 Location RT (124–138)

**Author Location** Pol (303–317)

Epitope FRKYTAFTIPSINNE

**Epitope name** Pol 303

Immunogen vaccine

Vector/Type: DNA with CMV promotor, peptide Adjuvant: Complete Freund's Adjuvant

(CFA)

Species (MHC) mouse (I-Ab and HLA-DR)

Donor MHC H-2b

Keywords vaccine-specific epitope characteristics, im-

munodominance

References Livingston et al. 2002

- 4 Th HIV epitopes presented by HLA-DR molecules were identified that also could be presented by murine class II molecule I-Ab, enabling testing of vaccine strategies in H-2b mice.
- Responses to pooled peptides, polyepitope peptides in a linear construct or in a branched MAP construct, and a DNA polyepitope construct with a CMV promoter were compared. A linear

arrangement in polyepitope construct created a junctional epitope that could be disrupted with the addition of GPGPG spacers. The linear polyepitope construct with the GPGPG spacer worked well in terms of eliciting responses to all 4 peptides, using either DNA or protein for the vaccination.

HXB2 Location RT (133–147) Author Location RT (133–147 BRU) Epitope PSINNETPGIRYQYN

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: RT

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>i5</sup>) **References** De Groot *et al.* 1991

• T-cells from RT immunized mice have enhanced proliferative response with peptide.

HXB2 Location RT (144–158) Author Location RT (144–158 BRU) Epitope YQYNVLPQGWKGSPA

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: RT

**Species (MHC)** mouse (H-2<sup>t4</sup>) **References** De Groot *et al.* 1991

 T-cells from RT immunized mice have enhanced proliferative response with peptide.

HXB2 Location RT (156–170) Author Location Pol (335–349) Epitope SPAIFQSSMTKILEP

**Epitope name** Pol 596

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

**Keywords** subtype comparisons **References** Wilson *et al.* 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.
- This epitope binds nine HLA-DR alleles: DRB1\*0101, DRB1\*1501, DRB1\*0405, DRB1\*1101, DRB1\*1302, DRB1\*0701, DRB1\*0901, DRB5\*0101 and DRB3\*0101, with an IC $_{50}$  threshold below 1,000 nM.
- This epitope sequence is conserved in 79% of clade B isolates.
- 7/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 non-responder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

**HXB2 Location** RT (156–170) **Author Location** RT (335–349)

Epitope SPAIFQSSMTKILEP

Epitope name Pol2

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Country United Kingdom.

Assay type proliferation, Intracellular cytokine staining

**Keywords** supertype, rate of progression

References Boaz et al. 2003

- Proliferative and cytokine (IFNgamma and II-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+ T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+ T cell declined to <500 after 15 years. Patients were treatment-naive.</li>
- Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs.
   Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.
- Pol2 was 1 of 3 peptides that had a negative correlation between absolute number of responding cells and viral load.

HXB2 Location RT (156–170) Author Location Pol (335–449) Epitope SPAIFQSSMTKILEP

Epitope name Pol 335 Immunogen vaccine

Vector/Type: DNA with CMV promotor, peptide Adjuvant: Complete Freund's Adjuvant

(CFA)

Species (MHC) mouse (I-Ab and HLA-DR)

Donor MHC H-2b

**Keywords** vaccine-specific epitope characteristics, immunodominance

References Livingston et al. 2002

- 4 Th HIV epitopes presented by HLA-DR molecules were identified that also could be presented by murine class II molecule I-Ab, enabling testing of vaccine strategies in H-2b mice.
- Responses to pooled peptides, polyepitope peptides in a linear construct or in a branched MAP construct, and a DNA polyepitope construct with a CMV promoter were compared. A linear arrangement in polyepitope construct created a junctional epitope that could be disrupted with the addition of GPGPG spacers. The linear polyepitope construct with the GPGPG spacer worked well in terms of eliciting responses to all 4 peptides, using either DNA or protein for the vaccination.

**HXB2 Location** RT (171–189)

Author Location Pol (171–189 HXB2)

Epitope FRKQNPDIVIYQYMDDLYV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DR0101)

Assay type Cytokine production, proliferation, Tetramer binding, CD4 T-cell Elispot - IFN $\gamma$ 

**Keywords** HAART, ART, supervised treatment interrup-

tions (STI)

References Iyasere et al. 2003

Fifteen patients recieving HAART with strong CD4+ proliferative responses to HIV antigens while on therapy were examined, to see the effects of viremia on these responses during treatment interruptions. Increased viremia occurred in 12/15 patients during at least one treatment interruption. Anti-HIV proliferative responses were inhibited during viremia, but IFNgamma production to Gag, Pol, and Nef peptide pools were maintained.

 IL-2 production diminished during viremia, and exogenous IL-2 revived *in vitro* proliferation of HIV-specific T-cells to Gag or Pol DR0101 epitopes in a tetramer, as well as Gag-specific total CD4 T-cell responses.

**HXB2 Location** RT (171–190)

**Author Location** RT (171–190 HXB2)

Epitope FRKQNPDIVIYQYMDDLYVG

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DR1, DR2 or DR3, DR4 and DR7)

Keywords Th1

References van der Burg et al. 1999

- T-cells specific for this epitope from the three donors were stimulated when presented with target cells pulsed with whole RT, indicating that the peptide is naturally processed for multiple HLA-DR molecules.
- Epitope binds to HLA-DR1, -DR2, -DR3, -DR4, and DR7, and can elicit Th1 cells that recognize peptide, protein, and HIV pulsed stimulator cells in the context of DR1, 2 or 3, 4 and 7 these HLA types cover more than half of the general population.

**HXB2 Location** RT (171–190)

**Author Location** RT (171–190 HXB2)

Epitope FRKQNPDIVIYQYMDDLYVG

Subtype B

Immunogen HIV-1 infection, in vitro stimulation or selec-

tion

Species (MHC) human (DR1, DR2, DR3, DR4, DR7)

**Keywords** binding affinity, cross-presentation by different HLA. Th1

References van der Burg et al. 1999

- The goal of this study was to identify Th epitopes that could be cross-presented by multiple class II HLA molecules. 5 RT peptides were identified that could bind to more than one HLA class II protein, and but only 2/5 could stimulate strong proliferation responses in PBMC derived from multiple healthy donors.
- This highly conserved epitope binds with high affinity to HLA-DR1, -DR2, -DR3, -DR4, and -DR7 but not HLA-DR5, and stimulated proliferation in 3/3 PBMC individuals with the appropriate HLA alleles.
- This epitope was able to be naturally processed in protein pulsed stimulator cells, and responding clones had a Th1 cytokine profile.
- This epitope is highly conserved and spans the highly conserved YMDD motif, and showing only minor variability in clades A, B, and D.

**HXB2 Location** RT (195–209)

**Author Location RT (IIIB)** 

Epitope IGQHRTKIEELRQHL

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

• Protein priming induced T-cells that recognize peptide.

HXB2 Location RT (196–215) Author Location RT (351–370) Epitope GQHRTKIEELRQHLLRWGLT

**Immunogen** in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995a

 Protein priming induced T-cells that recognize peptide, 4 clones from a single donor recognized this peptide.

**HXB2 Location** RT (249–263)

**Author Location** RT (249–263)

Epitope KDSWTVNDIQKLVGK

Epitope name RT2

Immunogen vaccine, in vitro stimulation or selection

Vector/Type: HIV-1 peptide in filamentous bacteriophage major coat protein HIV com-

ponent: RT

Species (MHC) human (DR5)

**Keywords** epitope processing

References De Berardinis et al. 2000

- Phage display of the CTL epitope, ILKEPVHGV coupled with T helper epitope KDSWTVNDIQKLVGK, elicited specific CTL responses in PBMC from HIV negative individuals and in vivo in immunization of HLA-A2 transgenic mice.
- Bacteriophage presentation of peptides is generally used for stimulation of antibodies, and this novel discovery of CTL epitope processing and presentation suggests new possibilities for these vectors.
- HIV-1 peptides were displayed in filamentous bacteriophage fd virion major coat protein pVIII.

**HXB2 Location** RT (249–263)

Author Location RT (249-263)

Epitope KDSWTVNDIQKLVGK

Epitope name pep23

Immunogen vaccine, in vitro stimulation or selection

Vector/Type: peptide presented on icosahedral protein scaffold HIV component: RT Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human, transgenic mouse (DR5)

**Assay type** Cytokine production, T-cell Elispot, Th support of CTL response

**References** Domingo et al. 2003

- A virus-like protein scaffold, called E2DISP, derived from pyruvate dehydrogenase multienzyme from Bacillus stearothermophilus has been engineered to display 60 copies of one or more epitopes on a single molecule. An E2DISP scaffold which displayed pep23, a 15-residue B and T helper epitope from the reverse transcriptase of HIV-1 elicited a T-helper response in vitro.
- The E2DISP scaffold displaying pep23 to stimulate a Th responses, and peptide RT2, which is a CTL epitope from HIV-1 reverse transcriptase, was able to elicit a CD8+ T cell response *in vitro* and in a vaccinated HLA-A2 transgenic mouse. Thus the E2DISP scaffold allows cell-entry and access to the class I and class II processing pathways.
- The Th response in vaccinated mice was also able to support Pep23 specific IgG response.

HXB2 Location RT (249–263)

**Author Location** RT (248–262)

Epitope KDSWTVNDIQKLVGK

**Immunogen** in vitro stimulation or selection

Species (MHC) human (DR5-11.01)

**Donor MHC** DR5, DR6 **Assay type** proliferation

**Keywords** binding affinity, epitope processing, vaccinespecific epitope characteristics, escape

References Moschella et al. 2003

- Two helper T-cell clones specific for this epitope presented in the context of HLA-DR5-11.01 have been characterized. They have different T cell receptor usage. Residue 11 (kdswtvndiqK-lvgk)is a natural variant, and K11A, K11G, K11I, and K11L variants were synthesized and studied in two presentation contexts, one as simple peptides, the other embedded in a recombinant protein, GST.
- The two Th clones and the two presentation contexts gave different outcomes with the peptides. K11I was not stimulatory, and was an antagonist in GST, an agonist as a peptide. K11L retained reactivity when presented in the fusion antigen, and had no acitivity as a peptide. K11G stimulated in both contexts, but the concentrations required for half maximal reactivity were different. K11A could not bind to the MHC in the processed form and could only stimulate when given as a peptide.
- In conclusion, substitutions in epitopes have different effects on Th stimulation depending on the mode of processing, and this should be considered when interpreting Th escape studies and vaccine development.

**HXB2 Location** RT (249–263)

Author Location RT (248-262)

Epitope KDSWTVNDIQKLVGK

Immunogen in vitro stimulation or selection

Species (MHC) human (DR5-11.01)

Donor MHC DR5, DR6

Assay type proliferation

Keywords binding affinity, epitope processing, vaccine-

specific epitope characteristics, escape, TCR

usage

References Bonomi et al. 2000

- Two helper T-cell clones specific for this epitope presented in the context of HLA-DR5-11.01 have been characterized. One of them used TCR Vβ15, the other used Vβ2. The substituions D2A, W4A, D8A, I9A, and K15A were generated and only D8A, I9A failed to react with one clone, while W4A, D8A, I9A were all critical for a reaction with the other clone, showing the TCRs focused on different but overlapping residues.
- Moving the epitope to different contexts in recombinant proteins for presentation by APCs, as well as adding polyanalanine and polyserine strings to either side of the epitope, influenced reactivity, suggesting processing context can influence the structure of the presentation complex.

**HXB2 Location** RT (249–263)

**Author Location** RT (248–262 HXB2)

Epitope KDSSTVNDIQKLVGK

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DRS)

**References** Fenoglio *et al.* 1999

- RT pep23 epitope exhibited antagonistic activity against proliferation of gp120-specific T-cells when flanked by unrelated amino acid sequence.
- The glutathione S-transferase (GST)-peptide system can be used to display peptides; antigenicity was maintained when this peptide was expressed at the C-term end, but antagonism resulted when this peptide was expressed at the N-term end.

**HXB2 Location** RT (249–263)

**Author Location** (243–263)

Epitope KDSWTVNDIQKLVGK

Epitope name pep23 Immunogen vaccine

> Vector/Type: bacteriophage coat protein, dihydrolipoyl acetyltransferase E2 protein, of Bacillus stearothermophilus HIV compo-

nent: RT

Species (MHC) transgenic mouse (HLA-DR)

**Assay type** Chromium-release assay **Keywords** vaccine antigen design **References** De Berardinis *et al.* 2003

An RT T-helper (KDSWTVNDIQKLVGK) that can be promiscuously presented by multiple HLA-DR molecules, and an RT CTL epitope (ILKEPVHGV) presented by HLA-A2, were displayed using two different antigen presentation systems, bacteriophage virions or E2 protein scaffolds. Both systems enabled display of the epitopes in a mouse model system to the immune system. CTL responses were detected in immunized mice, and were processed correctly for both class I and class II presentation.

**HXB2 Location** RT (249-263)

Author Location RT (IIIB)

Epitope KDSWTWNDIQKLVGK

**Immunogen** in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro.
- Peptide priming did not induce T-cells that recognize whole protein.

**HXB2 Location** RT (249–263)

Author Location RT (248-262)

Epitope KDSWTVNDIQKLVGK

Immunogen in vitro stimulation or selection

Species (MHC) human

**References** De Berardinis *et al.* 1999

- PBMC from donors GD (HLA DR 11; DRB52) and LD (HLA DR 11, 13; DRB52) recognized this epitope (pep23)
- A subset of T-cell lines generated from these donors were capable of recognizing pep23 expressed on the surface of filamentous phage fd, fused to the major coat protein gVIIIp.
- This peptide was selected to study phage presentation of peptide sequences because it was known to serve as a T-cell helper determinant which could induce proliferation from a naive repertoire Manca *et al.* [1995a]

**HXB2 Location** RT (251–261)

**Author Location** RT (250–260)

Epitope SSTVNDIQKLV

Immunogen in vitro stimulation or selection

Species (MHC) human (DR5(11.01))

References Manca et al. 1996

- This peptide was the minimal stimulatory sequence.
- One Th line was stimulated by p66, one by a Glutathione-Stransferase (GST)-peptide fusion protein.
- Constructs linking GST to the KDSSTVNDIQKLVGK peptide at the N-term end of GST stimulated Th cells, but not constructs linking at the C-term end.
- The C and N termini of GST are not intrinsically permissive or non-permissive, presentation is epitope specific (see FAILKC-NNK for contrast)

**HXB2 Location** RT (258–272)

Author Location RT (IIIB)

Epitope QKLWGKLNWASQIYP

**Immunogen** in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro
- Peptide priming did not induce T-cells that recognize whole protein.

**HXB2 Location** RT (271–290)

**Author Location** RT (271–290 HXB2)

Epitope YPGIKVRQLCKLLRGTKALT

Subtype B

Immunogen HIV-1 infection, in vitro stimulation or selec-

ion

Species (MHC) human (DR1, DR2, DR3, DR5, DR7)

**Keywords** binding affinity, cross-presentation by different HLA

References van der Burg et al. 1999

- The goal of this study was to identify Th epitopes that could be cross-presented by multiple class II HLA molecules. 5 RT peptides were identified that could bind to more than one HLA class II protein, but only 2/5 could stimulate strong proliferation responses in PBMC derived from multiple healthy donors.
- This epitope binds with high affinity to HLA-DR1, -DR2, -DR3, -DR5, and -DR7 but not HLA-DR4, and stimulated proliferation in 3/4 individuals with the appropriate HLA alleles.
- This epitope was not able to be naturally processed in proteinpulsed stimulator cells.

**HXB2 Location** RT (271–290)

Author Location RT (271–290 HXB2)

Epitope YPGIKVRQLCKLLRGTKALT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References van der Burg et al. 1999

 Epitope can bind to at least 5 different HLA-DR molecules, and peptide on target cells can elicit Th responses from PBMC cultures from healthy donors, however it does not seem to be processed properly from whole RT or virus.

**HXB2 Location** RT (276–290)

**Author Location RT (IIIB)** 

Epitope WRQLCKLLRGTKALT

**Immunogen** in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

• Protein priming induced T-cells that recognize peptide.

**HXB2 Location** RT (285–299)

Author Location RT (IIIB)

**Epitope** GTKALTEVIPLTEEA

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

• Protein priming induced T-cells that recognize peptide.

**HXB2 Location** RT (294–308)

Author Location RT (IIIB)

**Epitope** PLTEEAELELAENRE

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

• Protein priming induced T-cells that recognize peptide.

**HXB2 Location** RT (303-317)

**Author Location** RT (IIIB)

Epitope LAENREILKEPVHGV

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

• Protein priming induced T-cells that recognize peptide.

**HXB2 Location** RT (384–398)

**Author Location** RT (IIIB)

Epitope GKTPKFKLPIQKETW

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

• Protein priming induced T-cells that recognize peptide.

**HXB2 Location** RT (414–428)

**Author Location** Pol (596–610)

Epitope WEFVNTPPLVKLWYQ

Epitope name Pol 596

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

**Keywords** subtype comparisons

References Wilson et al. 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.
- This epitope binds eleven HLA-DR alleles: DRB1\*0101, DRB1\*1501, DRB1\*0401, DRB1\*0405, DRB1\*1101, DRB1\*1302, DRB1\*0701, DRB1\*0802, DRB1\*0901, DRB5\*0101 and DRB4\*0101, with an IC<sub>50</sub> threshold below 1,000 nM.
- This epitope sequence is conserved in 84% of clade B isolates.
- 6/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 non-responder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

**HXB2 Location** RT (414–428) **Author Location** RT (596–610)

Epitope WEFVNTPPLVKLWYQ

Epitope name Pol3

Immunogen HIV-1 infection

**Species (MHC)** human (DR(supermotif))

**Country** United Kingdom.

**Assay type** Cytokine production, proliferation **Keywords** supertype, rate of progression

References Boaz et al. 2003

- Proliferative and cytokine (IFNgamma and Il-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+ T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+ T cell declined to <500 after 15 years. Patients were treatmentnaive</li>
- Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs.
   Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.
- Pol3 was 1 of 2 peptides that had a positive correlation between absolute number and percentage of responding cells and viral load. Pol3 responses were also negatively correlated with CD4 counts. In contrast, the absolute number of 3/11 peptides studied were negatively correlated with viral load.

**HXB2 Location** RT (429–443)

**Author Location** RT (IIIB)

Epitope LEKEPIVGAETFYVD

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

• Protein priming induced T-cells that recognize peptide.

**HXB2 Location** RT (432–450)

Author Location RT (431–450 HXB2)

Epitope EPIVGAETFYVDGAANRET

Subtype B

Immunogen HIV-1 infection, in vitro stimulation or selec-

Species (MHC) human (DR1, DR2, DR3, DR4)

**Keywords** binding affinity, cross-presentation by different HLA

References van der Burg et al. 1999

- The goal of this study was to identify Th epitopes that could be cross-presented by multiple class II HLA molecules. 5 RT peptides were identified that could bind to more than one HLA class II protein, and but only 2/5 could stimulate strong proliferation responses in PBMC derived from multiple healthy donors.
- This epitope binds with high affinity to HLA-DR1, -DR2, -DR3, and -DR4, but stimulated a strong proliferation response in only 1/4 individuals tested so was not considered broadly cross-presented.

**HXB2 Location** RT (526–540)

**Author Location** RT (526–540 BRU)

Epitope IKKEKVYLAWVPAHK

Epitope name W9 Subtype B

Immunogen vaccine

Vector/Type: peptide, protein, inactivated HIV Strain: B clade BRU HIV component: RT, virus Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (Ad, Dd)

References Haas et al. 1991

- Of 5 mouse inbred lines tested: DBA/2 (H-2d, Ad, Ed), B10.A(4R) (H-2h4, Ak) and B10.A(5R) (H-2i5) showed particularly good CD4+ T cell proliferative responses to HIV proteins (gp160, gp120, p17, p24, Nef and RT), after vaccination with inactivated virus.
- B10.BR (H-2k, Ak, Ek) and C57BL/6 (H-2b and Ab) had weaker responses.
- The peptide KEKVYLAWVPAHKG was one of two RT peptides with Th cells recognition.

**HXB2 Location** RT (528–540)

Author Location RT (528-540)

Epitope KEKVYLAWVPAHK

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade BRU HIV component: RT Adjuvant: P3CSS

Species (MHC) mouse (H-2b, H-2d, H-2k)

**Assay type** proliferation **References** Loleit *et al.* 1996

• BALB/c, C3H/Hej, and C57BL/6 mice were immunized with 22-mer lipopetide tripeptide conjugates P3CSS-[RT-(522-543)] and P3CSS-[RT-(528-549)] of HIV-1 RT, which included the optimal T-helper epitope [RT-(528-540)]. P3CSS conjugated RT epitopes resulted in a specific Th responses, and mice were primed for secondary recognition of native RT. A proximal B cell epitope was also active, containing the motif EQVD.

**HXB2 Location** RT (528–541)

**Author Location** RT (528–543 BRU)

Epitope KEKVYLAWVPAHKG

Epitope name A3

Subtype B

Immunogen vaccine

Vector/Type: peptide, protein, inactivated HIV Strain: B clade BRU HIV component: RT, virus Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (Ad, Dd)

Donor MHC H-2d, H-2f, H-2k

References Haas et al. 1991

- Of 5 mouse inbred lines tested: DBA/2 (H-2d, Ad, Ed), B10.A(4R) (H-2h4, Ak) and B10.A(5R) (H-2i5) showed particularly good CD4+ T cell proliferative responses to HIV proteins (gp160, gp120, p17, p24, Nef and RT), after vaccination with inactivated virus.
- B10.BR (H-2k, Ak, Ek) and C57BL/6 (H-2b and Ab) had weaker responses.

 The peptide KEKVYLAWVPAHKG was one of two RT peptides with Th cells recognition. It could by itself prime different strains of mice for RT-specific Th responses, and the C-term half of the peptide is highly conserved in HIV-1, HIV-2 and SIV strains.

HXB2 Location RT (528–543) Author Location RT (528–543 BRU) Epitope KEKVYLAWVPAHKGIG

Immunogen vaccine

Vector/Type: peptide Strain: B clade BRU

**Species (MHC)** mouse  $(H-2^f, H-2^k, H-2^d)$ 

References Haas et al. 1991

 T-cells from peptide-primed mice could be restimulated by native RT.

Author Location RT (529–543)

Author Location Pol (711–725)

Epitope EKVYLAWVPAHKGIG

Epitope name Pol 711

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif) Keywords subtype comparisons References Wilson *et al.* 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.
- This epitope binds ten HLA-DR alleles: DRB1\*0101, DRB1\*1501, DRB1\*0401, DRB1\*0405, DRB1\*1101, DRB1\*0701, DRB1\*0802, DRB1\*0901, DRB5\*0101 and DRB4\*0101, with an IC<sub>50</sub> threshold below 1,000 nM.
- This epitope sequence is conserved in 94% of clade B isolates.
- 6/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 non-responder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

**HXB2 Location** RT (529–543)

Author Location Protease-RT (711–725)

Epitope EKVYLAWVPAHKGIG

Epitope name Pol4

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Assay type proliferation, Intracellular cytokine staining

Keywords rate of progression, superinfection

References Boaz et al. 2003

- Proliferative and cytokine (IFNgamma and II-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+ T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+ T cell declined to <500 after 15 years. Patients were treatment-naive</li>
- Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs.
   Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated

with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.

HXB2 Location RT (529–543) Author Location Pol (711–725) Epitope EKVYLAWVPAHKGIG

Epitope name Pol 711 Immunogen vaccine

Vector/Type: DNA with CMV promotor, peptide Adjuvant: Complete Freund's Adjuvant

(CFA)

Species (MHC) mouse (I-Ab and HLA-DR)

Donor MHC H-2b

**Keywords** vaccine-specific epitope characteristics, immunodominance

References Livingston et al. 2002

- 4 Th HIV epitopes presented by HLA-DR molecules were identified that also could be presented by murine class II molecule I-Ab, enabling testing of vaccine strategies in H-2b mice.
- Responses to pooled peptides, polyepitope peptides in a linear construct or in a branched MAP construct, and a DNA polyepitope construct with a CMV promoter were compared. A linear arrangement in polyepitope construct created a junctional epitope that could be disrupted with the addition of GPGPG spacers. The linear polyepitope construct with the GPGPG spacer worked well in terms of eliciting responses to all 4 peptides, using either DNA or protein for the vaccination.
- Although responses to this peptide indicated it was immunodominant, responses to all 4 peptides were made upon vaccination with linear constructs when GPGPG spacers were used.

**HXB2 Location** RT (530–544) **Author Location** Pol (712–726)

Epitope KVYLAWVPAHKGIGG

Epitope name Pol 712

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Keywords subtype comparisons

References Wilson et al. 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.
- This epitope binds ten HLA-DR alleles: DRB1\*0101, DRB1\*1501, DRB1\*0401, DRB1\*0405, DRB1\*1101, DRB1\*0701, DRB1\*0802, DRB1\*0901, DRB5\*0101 and DRB4\*0101, with an IC $_{50}$  threshold below 1,000 nM.
- This epitope sequence is conserved in 89% of clade B isolates.
- 6/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 non-responder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

**HXB2 Location** RT (530–544) **Author Location** RT (712–726)

Epitope KVYLAWVPAHKGIGG

Epitope name Pol5

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Country United Kingdom.

Assay type proliferation, Intracellular cytokine staining

Keywords supertype, rate of progression

References Boaz et al. 2003

- Proliferative and cytokine (IFNgamma and II-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+T cell declined to <500 after 15 years. Patients were treatment-naive.</li>
- Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs.
   Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.
- Pol5 was 1 of 2 peptides that had a positive correlation between absolute number and percentage of responding cells and viral load. In contrast, the absolute number of 3/11 peptides studied were negatively correlated with viral load.

# III-B-6 RT-Integrase Helper/CD4 + T-cell epitopes

**HXB2 Location** RT-Integrase (553–3)

**Author Location** RT (720–730 LAI)

Epitope SAGIRKVLFLD

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

# III-B-7 Integrase Helper/CD4+ T-cell epitopes

**HXB2 Location** Integrase (16–30)

Author Location Pol (758-772)

Epitope HSNWRAMASDFNLPP

Epitope name Pol 758

Immunogen HIV-1 infection

**Species** (MHC) human (DR supermotif)

**Keywords** subtype comparisons

References Wilson et al. 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.
- This epitope binds eight HLA-DR alleles: DRB4\*0101, DRB5\*0101, DRB1\*0901, DRB1\*0701, DRB1\*1101, DRB1\*0405, DRB1\*0401 and DRB1\*0101, with an IC<sub>50</sub> threshold below 1,000 nM.
- This epitope sequence is conserved in 68% of clade B isolates.

• 8/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 non-responder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

**HXB2 Location** Integrase (16–30)

Author Location Integrase (758-772)

Epitope HSNWRAMASDFNLPP

Epitope name Pol6

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Country United Kingdom.

Assay type proliferation, Intracellular cytokine staining

Keywords supertype, rate of progression

References Boaz et al. 2003

- Proliferative and cytokine (IFNgamma and II-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+T cell declined to <500 after 15 years. Patients were treatment-naive.</li>
- Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs.
   Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.
- Pol6 was 1 of 3 peptides that had a negative correlation between absolute number of responding cells and viral load.

HXB2 Location Integrase (172–186)

Author Location RT (899–913 LAI)

Epitope LKTAVQMAVFIHNFK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

**HXB2 Location** Integrase (173–187)

Author Location Pol (915-929)

Epitope KTAVQMAVFFIHNFKR

Epitope name Pol 915

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Keywords subtype comparisons

References Wilson et al. 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.
- This epitope binds seven HLA-DR alleles: DRB5\*0101, DRB1\*1302, DRB1\*1101, DRB1\*0405, DRB1\*0401, DRB1\*1501 and DRB1\*0101, with an IC<sub>50</sub> threshold below 1.000 nM
- This epitope sequence is conserved in 94% of clade B isolates.

responded to some of the DR supermotif epitopes, the 9 nonresponder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

HXB2 Location Integrase (173–187) Author Location Integrase (915–929) Epitope KTAVQMAVFIHNFKR

Epitope name Pol7

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Country United Kingdom.

Assay type proliferation, Intracellular cytokine staining Keywords supertype, rate of progression, immunoprophylaxis

References Boaz et al. 2003

- Proliferative and cytokine (IFNgamma and Il-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+ T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+ T cell declined to <500 after 15 years. Patients were treatment-
- Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs. Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.

HXB2 Location Integrase (196-210) **Author Location** RT (923–937 LAI)

Epitope AGERIVDIIATDIQT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

HXB2 Location Integrase (214–228)

Author Location Pol (956–970)

Epitope QKQITKIQNFRVYYR

Epitope name Pol 956

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

**Keywords** subtype comparisons

References Wilson et al. 2001

- Eleven peptides were identified that had the HLA-DR supermotif, all were found to bind to MHC class II DR molecules and all elicited proliferative responses from multiple HIV-infected donors.
- This epitope binds twelve HLA-DR alleles: DRB4\*0101, DRB5\*0101, DRB1\*0901, DRB1\*0802, DRB1\*0701, DRB1\*1302, DRB1\*1201, DRB1\*1101, DRB1\*0405, DRB1\*0401, DRB1\*1501 and DRB1\*0101, with an IC50 threshold below 1,000 nM.
- This epitope sequence is conserved in 95% of clade B isolates.

• 6/22 HIV infected individuals responded to this epitope (13/22 • 8/22 HIV infected individuals responded to this epitope (13/22 responded to some of the DR supermotif epitopes, the 9 nonresponder peptides tended to also not have recall responses to rec HIV-1 whole proteins)

HXB2 Location Integrase (214–228)

Author Location Integrase (956–970)

Epitope QKQITKIQNFRVYYR

Epitope name Pol8

Immunogen HIV-1 infection

Species (MHC) human (DR supermotif)

Country United Kingdom.

Assay type proliferation, Intracellular cytokine staining

Keywords supertype, rate of progression

References Boaz et al. 2003

- Proliferative and cytokine (IFNgamma and Il-2) immune responses to 11 previously defined (Wilson 2001), and immunodominant and broadly cross-reactive Th epitopes and to p24 overlapping peptides were characterized in 10 LTNPs (chronically infected, asymptomatic, long term non-progressors, CD4+ T cells >500 after 15 years) and 7 SPs (slow progressors, CD4+ T cell declined to <500 after 15 years. Patients were treatmentnaive.
- Gag-specific CD4+ T cells in LTNPs showed increased numbers of IFN gamma and IL-2 producing cells compared to SPs. Cytokine production and proliferative responses were negatively correlated with the viral load and positively correlated with the CD4+ T cell count. IL-2, but not IFNgamma, levels were correlated with proliferation.
- Pol8 was the only peptide that had higher cytokine responses in LTNPs than SPs (p = 0.0431). No peptide had detectable differences in proliferative responses between the two groups.

HXB2 Location Integrase (215-227)

**Author Location** RT (942–954 LAI)

Epitope KQITKIQNFRVYY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

HXB2 Location Integrase (250–267)

Author Location Integrase (250–267 B Consensus)

Epitope VIQDNSDIKVVPRRKAKI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

• CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.

- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location Integrase (250–267) Author Location Integrase (250–267)

Epitope VIQDNSDIKVVPRRKAKI

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production

References Geels et al. 2006

- The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.
- Autologous sequences corresponding to known and predicted Thepitopes were analyzed. VIQDNSDIKVVPRRKAKI had fixation of 1 mutation (VIQDNSDIK[v/a]VPRRKAKI) in 1 of the patients.

## III-B-8 Pol Helper/CD4+ T-cell epitopes

HXB2 Location Pol

**Author Location** RT (248–256 HXB2)

Epitope Subtype B

**Immunogen** in vitro stimulation or selection

Species (MHC) human (DR5)

References Manca et al. 1995b

- CD4+ T-cell lines from uninfected individuals by stimulation with p66-pulsed APC.
- TcR V $\beta$  D $\beta$  J $\beta$  sequences were obtained from p66-specific T-cell clones.
- There were multiple responses to peptides throughout p66, but because of uncertain locations, they have not been mapped.
- Response to peptide 248-256 was associated with DR5.

HXB2 Location Pol Author Location RT

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env, Gag, Pol Adjuvant: IFNγ, IL-2, IL-4

Species (MHC) mouse (H-2<sup>d</sup>)

Keywords Th1

References Kim et al. 2000

• Co-stimulatory molecules co-expressed with an HIV-1 immunogen in a DNA vaccine used to enhance the immune response – co-expression of Th1 cytokine IFN $\gamma$  drove Th1 immune responses and enhanced CTL responses.

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

Immunogen vaccine

Vector/Type: Salmonella HIV component:

RT

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Burnett et al. 2000

 A live attenuated bacterial vaccine, Salmonella SL3261pHART, with an inserted HIV RT gene in the Lpp-OmpA-HIV fusion protein, induced a lymphoproliferative Th response in BALB/c mice.

HXB2 Location Pol

Author Location Gag/Pol

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Gag,

Pol, Vif Adjuvant: B7, IL-12

Species (MHC) mouse

References Kim et al. 1997b

A gag/pol DNA vaccine delivered in conjunction with the plasmid encoding the co-stimulatory molecules B7 and IL-12 gives a dramatic increase in both the cytotoxic and proliferative responses in mice.

HXB2 Location Pol

Author Location Gag/Pol

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Gag,

gp160, Pol Adjuvant: CD86

Species (MHC) mouse

References Kim et al. 1997d

A gag/pol DNA vaccine delivered in conjunction with the plasmid encoding the co-stimulatory molecule CD86 gives an increase in proliferative responses to Pr55 in mice.

HXB2 Location Pol

Author Location Gag/Pol (MN)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade MN HIV component: Env, Gag, Pol Adjuvant:

CD80, CD86

Species (MHC) chimpanzee

References Kim et al. 1998

Co-stimulatory molecules co-expressed with an HIV-1 immunogen in a DNA vaccine used to enhance the immune response – co-expression of CD86, but not CD80, dramatically increased both HIV Env and Gag/Pol specific CTL and Th proliferative responses.

HXB2 Location Pol

Author Location Pol Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART **References** Blankson *et al.* 2001

- 5/10 chronically HIV infected patients with low CD4+ counts who received HAART therapy and experienced immune reconstitution displayed p24, p17 and p66 T-helper CD4 proliferative responses, in contrast to 0/8 chronically HIV infected patients with high CD4+ counts at the initiation of antiretroviral treatment.
- This surprising result could be due to the low CD4 nadir patients being more likely to have thymic regeneration or a peripheral expansion of T cells.

HXB2 Location Pol Author Location p66 Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART **References** Oxenius *et al.* 2000

 Patients who started therapy at acute HIV infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.

HXB2 Location Pol
Author Location p66
Epitope
Immunogen HIV-1 infection
Species (MHC) human
Keywords HAART, ART
References Palmer et al. 2002

- CD4+ T cell proliferative responses from 33 HIV-1 infected patients with HAART suppression were compared to 19 patients with active viral replication (HAART failures and HAART naive). Patients with HAART suppression showed stronger p24- and p66-specific proliferative responses compared to patient groups with active HIV-1 replication, suggesting active viral replication *in vivo* specifically reduces proliferation responses.
- No differences in the frequency of HIV-specific CD4+ T-cells that were positive for cytokine secretion in a flow cytometry assay were found in the HAART suppressed group versus the group with active viral replication.

HXB2 Location Pol Author Location (BRU) Epitope Subtype B Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade BRU HIV component: RT, virus Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse

References Haas et al. 1991

- Of 5 mouse inbred lines tested DBA/2 (H-2d, Ad, Ed), B10.A(4R) (H-2h4, Ak) and B10.A(5R) (H-2i5) showed particularly good CD4+ T cell proliferative responses to HIV proteins (gp160, gp120, p17, p24, Nef and RT), after vaccination with inactivated virus.
- B10.BR (H-2k, Ak, Ek) and C57BL/6 (H-2b and Ab) had weaker responses.

### III-B-9 Vif Helper/CD4 + T-cell epitopes

HXB2 Location Vif (65–76)

Author Location Vif (65–80)

Epitope VITTYWGLHTGE

Immunogen HIV-1 infection

Species (MHC) human

References Ranki et al. 1997

• T-cell response to this epitope persisted after seroreversion.

**HXB2 Location** Vif (81–96) **Author Location** Vif (81–96)

Epitope LGQGVSIEWRKQRYST Immunogen HIV-1 infection Species (MHC) human

References Ranki et al. 1997

• T-cell response to this epitope persisted after seroreversion.

HXB2 Location Vif Author Location Vif Epitope

Immunogen vaccine

Vector/Type: DNA HIV component: Nef, Vif, Vpu

**Species (MHC)** mouse (H-2<sup>d</sup>)

**Keywords** subtype comparisons, Th1 **References** Ayyavoo *et al.* 2000

- Splenocytes from BALB/c mice immunized with pVVN-P DNA were incubated with Vif, Vpu or Nef antigens for 3 days and assayed for IL-4 and IFNγ levels.
- Antigen stimulation increased IFN $\gamma$  production in pVVN-P immunized mice, indicating a Th1 response.
- IL-4 production was not significantly changed after antigen stimulation compared to control levels.
- Cross-clade CTL activity was also observed: A, B clade, CRF01(AE) clade antigens could serve as targets for the B clade immunization stimulated CTL an HIV-1 AC recombinant, however, did not stimulate a CTL response, but was expressed at lower levels on the target cell.

### III-B-10 Vpr Helper/CD4+ T-cell epitopes

**HXB2 Location** Vpr (66–80) **Author Location** Vpr (66–80 IIIB)

Epitope QLLFIHFRIGCRHSR

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Sarobe et al. 1994

• Included as a Th stimulatory component of peptide vaccines that also incorporated B-cell epitopes.

HXB2 Location Vpr (66–80)

Author Location Vpr (66–80 IIIB)

Epitope QLLFIHFRIGCRHSR

Immunogen HIV-1 infection

Species (MHC) human

References Sarobe et al. 1994

• This peptide was found to stimulate proliferative responses in 37.5% of HIV-1 positive individuals.

### III-B-11 Tat Helper/CD4 + T-cell epitopes

HXB2 Location Tat (1–20)

**Author Location** Tat (1–20 LAI)

Epitope MEPVDPRLEPWKHPGSQPKT

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

**HXB2 Location** Tat (16–35) **Author Location** Tat (16–35 LAI)

**Epitope** SQPKTACTTCYCKKCCFHCQ

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Tat (17–32)

**Author Location** Tat (17–32 HXB2)

Epitope QPKTACTNCYCKKCCF

Epitope name D26

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DR5? plus others)

Keywords immunodominance

References Blazevic et al. 1993

- 9/14 (64%) of HIV-1 positive patients had proliferative T-cell responses associated with IL-2 production against at least one Tat-derived synthetic peptides of twelve overlapping 15-16 mer peptides spanning Tat. T cell proliferation was associated with IL-2 production.
- 3/12 peptides were recognized.
- This immunodominant, highly conserved and most frequently recognized peptide was recognized by 57% of the HIV-1 infected patients. A beta-sheet secondary structure was predicted at aa residues 21-28, but no amphipathic helix structure, suggested to be most favorable for T-cell epitopes, was indicated.
- This peptide contained epitopes restricted by several HLA DR alleles, although the frequency of DR5 was enriched (3/6) among the patients that recognized the peptide.

HXB2 Location Tat (17–32)

**Author Location** Tat (17–32)

Epitope QPKTACTNCYCKRCCF

Immunogen HIV-1 infection

Species (MHC) human

References Ranki et al. 1997

• T-cell response to this epitope persisted after seroreversion.

**HXB2 Location** Tat (31–50)

**Author Location** Tat (31–50 LAI)

Epitope CFHCQVCFTTKALGISYGRK

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Tat (33–48)

**Author Location** Tat (33–48 HXB2)

Epitope HCQVCFITKALGISYG

Epitope name D28

Immunogen HIV-1 infection

Species (MHC) human (DR5? plus others)

References Blazevic et al. 1993

- 9/14 (64%) of HIV-1 positive patients had proliferative T-cell responses associated with IL-2 production against at least one Tat-derived synthetic peptides of twelve overlapping 15-16 mer peptides spanning Tat. T cell proliferation was associated with IL-2 production.
- 3/12 peptides were recognized.
- 4/14 HIV+ people recognized this peptide.
- An alpha-helix structure was predicted at residues 39-44, but charge patterns did not indicate it was an amphipathic helix, suggested to be most favorable for T-cell epitopes.

• This peptide contained epitopes restricted by several HLA DR alleles, although the frequency of DR5 was enriched (2/4) among the patients that recognized the peptide.

HXB2 Location Tat (33-48) Author Location Tat (33-48)

**Epitope** HCQVCFMTKGLGISYG

Immunogen HIV-1 infection

Species (MHC) human

References Ranki et al. 1997

• T-cell response to this epitope persisted after seroreversion.

HXB2 Location Tat (36–50)

Author Location Tat (36–50 HTLV IIIB)

Epitope VCFITKALGISYGRK?

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA), macrophage activating lipopeptide-2 (MALP-2)

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type Cytokine production, proliferation, T-cell Elispot, Th support of CTL response

Keywords Th1, Th2, mucosal immunity

References Borsutzky et al. 2003

- BALB/c mice that were vaccinated intranasally with Tat protein plus mucosal adjuvant macrophage-activating lipopeptide-2 (MALP-2) had increased proliferative, antibody, and IFNγ producing T-cell responses than did with Tat+IFA delivered by the i.p. route.
- IFA as adjuvant stimulated a Th2-dominant response pattern. and MALP-2 as adjuvant shifted to a Th1 response. Anti-Tat IgG1 dominated the Ab response with IFA, IgG2b dominated with MALP-2. In animals vaccinated with Tat+MALP-2, IFNγ and IL-2 were the most prominent cytokines, with some IL-6. In contrast, in mice vaccinated with Tat+IFA, IL-6 was the dominant cytokine. Secreted IL-4, IL-5 and IL-10 were below the detection limit in both cases.
- The strongest proliferation of spenocytes was observed was after re-stimulation with residues 36-50 and 56-70.

HXB2 Location Tat (41-50)

Author Location Tat (40–50 C consensus)

Epitope KGLGISYGRK?

Subtype C

Immunogen vaccine

Vector/Type: DNA Strain: C clade consensus HIV component: Tat Adjuvant: ubiquitin

Species (MHC) mouse

Donor MHC H-2d

Assay type proliferation, CD4 T-cell Elispot - IFNγ

Keywords Th1, vaccine antigen design References Ramakrishna et al. 2004

• BALB/c and C57BL/6 mice were intramuscularly immunized with a codon optimized HIV-1 C-consensus Tat DNA vaccine that was linked to ubiquitin to facilitate rapid processing. Ubiquitin and codon optimization enhanced Th1 T cell responses, with increased proliferative responses, cytotoxic responses, and

Th1 responses measured by IFNgamma EliSpot, but not the Th2 responses, measured by IL-4 EliSpot...

Several immunogenic regions in HIV-1 Tat were identified in BALB/c mice using EliSpot. The strongest immune response was within the core region of Tat; the peptides based on the C subtype consensus positions 30-50 and 40-60 gave the strongest EliSpot responses in BALB/c mice, suggesting a putative helper T-cell epitope spanning the region of overlap, residues 40-50.

HXB2 Location Tat (46–65)

**Author Location** Tat (46–65 LAI)

Epitope SYGRKKRRQRRRPPQGSQTH

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Tat (56–70)

**Author Location** Tat (56–70 HTLV IIIB)

Epitope RRAHQNSQTHQASLS?

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA), macrophage activating lipopeptide-2 (MALP-2)

**Species (MHC)** mouse (H-2<sup>d</sup>)

Assay type Cytokine production, proliferation, T-cell Elispot, Th support of CTL response

Keywords Th1, Th2

References Borsutzky et al. 2003

- BALB/c mice that were vaccinated intranasally with Tat protein plus mucosal adjuvant macrophage-activating lipopeptide-2 (MALP-2) had increased proliferative, antibody, and IFNγ producing T-cell responses than did with Tat+IFA delivered by the i.p. route.
- IFA as adjuvant stimulated a Th2-dominant response pattern, and MALP-2 as adjuvant shifted to a Th1 response. Anti-Tat IgG1 dominated the Ab response with IFA, IgG2b dominated with MALP-2. In animals vaccinated with Tat+MALP-2, IFNγ and IL-2 were the most prominent cytokines, with some IL-6. In contrast, in mice vaccinated with Tat+IFA, IL-6 was the dominant cytokine. Secreted IL-4, IL-5 and IL-10 were below the detection limit in both cases.
- The strongest proliferation of spenocytes was observed was after re-stimulation with residues 36-50 and 56-70.

HXB2 Location Tat (61–80)

**Author Location** Tat (61–80 LAI)

Epitope GSQTHQVSLSKQPTSQPRGD

Subtype B

Immunogen vaccine

HIV component: Nef. Rev. Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- · Stronger, broader responses were observed in animals vaccinated with DNA epidermally; rather than with intramuscular
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Tat (65-80)

**Author Location** Tat (65–80 HXB2)

Epitope HQASLSKQPTSQPRGD

Epitope name D32 Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DR2? plus others)

References Blazevic et al. 1993

- 9/14 (64%) of HIV-1 positive patients had proliferative T-cell responses associated with IL-2 production against at least one Tat-derived synthetic peptides of twelve overlapping 15-16 mer peptides spanning Tat. T cell proliferation was associated with IL-2 production.
- 3/12 Tat peptides were recognized.
- 3/14 HIV+ people recognized this peptide.
- An alpha-helix structure was predicted at residues 65-72, but charge patterns did not indicate it was an amphipathic helix, suggested to be most favorable for T-cell epitopes..
- This peptide contained epitopes restricted by several HLA DR alleles, although the frequency of DR2 was enriched (2/3) among the patients that recognized the peptide.

HXB2 Location Tat (67-86)

Author Location Tat (67–86 LAI)

Epitope VSLSKQPTSQPRGDPTGPKE

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- · Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

**HXB2 Location** Tat

**Author Location** Tat

**Epitope** 

Immunogen vaccine

Vector/Type: DNA, DNA with protein boost Strain: B clade LAI HIV component: Gag,

Nef, Tat Adjuvant: IL-18

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1, Th2

References Billaut-Mulot et al. 2001

• DNA vaccinated BALB/c mice primed and boosted with a multiepitopic vaccine with IL18 gave lymphoproliferative responses 7 weeks post immunization.

- Vector/Type: DNA Strain: B clade LAI Strong but non-lasting HIV-specific CTL responses were detected by a Cr-release assay and DNA prime + DNA boost was more effective than DNA prime + protein boost.
  - Immunization with either the multiepitopic DNA or with the mixed DNA vaccine resulted in Th1 cytokines production (IL-2 and IFN $\gamma$ ) in spleen cell cultures stimulated by Tat and Gag, while Th2 cytokines IL-4 and IL-10 production was not detect-
  - Co-administration of IL18 increased T-cell responses but decreased anti-HIV antibody levels.

**HXB2 Location** Tat

**Author Location** Tat

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Nef,

Rev, Tat

Species (MHC) human

Keywords HAART, ART

References Calarota et al. 1999

- 9/9 HIV-1 + subjects were given one of three DNA vaccinations for nef, rev or tat, and novel proliferative and CTL responses were generated.
- The nef DNA immunization induced the highest and most consistent CTLp activity, IFN \u03c4 production, and IL-6 and IgG
- Highly active antiretroviral treatment (HAART) did not induce new HIV-specific CTL responses but reduced viral load, while DNA vaccination induced new immune responses but did not reduce viral load – thus this is a potentially complementary and promising combination.

HXB2 Location Tat

**Author Location** Tat

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA HIV component: Nef, Rev, Tat Adjuvant: CpG immunostimula-

tory sequence (ISS)

Species (MHC) human

Keywords review, Th1

References Calarota & Wahren 2001

• This review discusses the cellular immune response, and comments on CpG induction of Th1 cytokines and enhanced immune responses, and HIV-1 DNA vaccine boosting of CTL and Th proliferative responses in asymptomatic HIV+ individuals.

**HXB2 Location** Tat

**Author Location** Tat

**Epitope** 

Immunogen in vitro stimulation or selection

Species (MHC) human

Keywords dendritic cells, Th1, Th2

References Corinti et al. 2002

• In vitro delivery of recombinant Tat protein conjugated to red blood cells (RBCs) via avidin-biotin bridges (RBC-Tat) to human dendritic cells was compared to dendritic cells pulsed with rec Tat.

- Dendritic cells pulsed with RBC-Tat elicited specific and significantly stronger CD4+ and CD8+ T-cell responses and required 1250-fold less antigen than DCs stimulated with soluble Tat.
- · Dendritic cells which were maturated in the presence of IFNgamma induced elevated IL-12 and TNF-alpha secretion. IFNgamma upregulated IP-10 and down regulated TARC, chemokines which attract Th1 and Th2 cells, respectively.

**HXB2 Location** Tat

Author Location Tat (IIIB, BH10)

**Epitope** Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human

Keywords epitope processing, vaccine-specific epitope characteristics, dendritic cells, Th1

References Fanales-Belasio et al. 2002b

- Biologically active HIV-1 Tat is readily taken up by monocytederived dendritic cells (MDDC) (and activated endothelial cells), but not other APCs. Tat must be in a native, non-oxidized conformation for efficient uptake. Tat upregulates MHC molecules, IL-12, TNF $\alpha$ , RANTES and MIP-1- $\alpha$  and MIP-1- $\beta$ production which drives Th1 immune responses and enhances antigen presentation.
- Native Tat enhanced the antigen presentation of MDDC and boosted proliferative recall and allogeneic antigen responses, and the authors propose it could be used as an adjuvant to drive the immune response as well as an antigen.

**HXB2 Location** Tat **Author Location** Tat

**Epitope** 

Immunogen vaccine

Vector/Type: DNA, protein HIV component: Tat Adjuvant: aluminum hydroxide, Ribi adjuvant (MPL+TDM) (RIBI)

Species (MHC) macaque

Assav type Cytokine production, Delayed-type hypersensitivity (DTH)

Keywords review, early-expressed proteins, Th1 References Fanales-Belasio et al. 2002a

• HIV-1 Tat protein has several virtues vaccine component. It is an early expressed protein, and though variable, contains consverved T-cell and B-cell epitopes that allow cross-clade recognition. It is efficiently taken up by monocyte-derived dendritic cells (MDDCs) and in this context can stimulate Th1 immune responses. A Tat based vaccine can elicit an immune response that can control primary infection in monkeys that are in early stage of in infection with SHIV89.6P.

**HXB2 Location** Tat

**Author Location** Tat (1–72)

**Epitope** Subtype B Immunogen vaccine

> Vector/Type: protein, nanoparticle Strain: B clade BRU HIV component: Tat Adjuvant:

aluminum hydroxide, lipid A

Species (MHC) mouse Donor MHC H-2d

Assay type Cytokine production, proliferation

Keywords Th1, Th2, adjuvant comparison, vaccine antigen design

References Cui et al. 2004

- Mice were subcutaneously injected on day 0 and 14 with either Alum and Tat (Th2 control) or Lipid A-adjuvanted Tat (Th1 control), or Tat coated anionic nanoparticles. Analysis of Ab and cytokine release in splenocytes (day 28) showed both IgG and IgM Ab responses; immunization with Tat-coated nanoparticles induced a Th1-biased immune response.
- IFN gamma responses were 3.3-fold stronger with Tat and either Lipid-A or coated nanoparticles than with Tat and Alum.

## III-B-12 Rev Helper/CD4 + T-cell epitopes

HXB2 Location Rev (9-23)

Author Location Rev (9-23 HXB2)

Epitope DEELIRTVRLIKLLY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Blazevic et al. 1995

• One of four peptides that stimulates in PBLs from HIV-1 + donors both CD4+ Th cell proliferation and CTL to autologous targets incubated with peptide were stimulated.

HXB2 Location Rev (14–30)

**Author Location** Rev (14–30 B Consensus)

**Epitope KTVRLIKFLYQSNPPPS** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNy EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location Rev (16-35)

**Author Location** Rev (16–35 LAI)

Epitope VRLIKFLYQSNPPPNPEGTR

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species** (MHC) mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Rev (25-39)

**Author Location** Rev (25–39 HXB2)

Epitope SNPPPNPEGTRQARR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Blazevic et al. 1995

 One of four peptides that stimulates in PBLs from HIV-1 + donors both CD4+ Th cell proliferation and CTL to autologous targets incubated with peptide were stimulated.

HXB2 Location Rev (31–50)

**Author Location** Rev (31–50 LAI)

Epitope PEGTRQARRNRRRRWRERQR

**Subtype** B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

**References** Hinkula *et al.* 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Rev (33-48)

**Author Location** Rev (33–48 HXB2)

Epitope GTRQARRNRRRRWRER

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Blazevic et al. 1995

• One of four peptides that stimulates in PBLs from HIV-1 + donors both CD4+ Th cell proliferation and CTL to autologous targets incubated with peptide were stimulated.

HXB2 Location Rev (41–56)

**Author Location** Rev (41–56 HXB2)

Epitope RRRRWRERQRQIHSIS

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Blazevic et al. 1995

 One of four peptides that stimulates in PBLs from HIV-1 + donors both CD4+ Th cell proliferation and CTL to autologous targets incubated with peptide were stimulated.

HXB2 Location Rev (76-95)

**Author Location** Rev (76–95 LAI)

Epitope PPLERLTLDCNEDCGTSGTQ

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Rev (96-116)

**Author Location** Rev (96–116 LAI)

Epitope GVGSPQILVESPTVLESGTKE

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

**HXB2 Location** Rev

**Author Location** Rev

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Rev

Species (MHC) mouse

Keywords HAART, ART

References Chan et al. 1998

- Rev M10 is a construct that was introduced into mice through a genetic vaccination.
- Rev was used to test for down-regulation of HIV-1 in infected cells as a method for gene therapy in the course of this study, Rev-specific IL-2 producing Th cells developed in the mice.

**HXB2 Location** Rev

**Author Location** Rev

Epitope

Immunogen vaccine

Vector/Type: DNA HIV component: Nef,

Rev. Tat

Species (MHC) human

Keywords HAART, ART

References Calarota et al. 1999

• 9/9 HIV-1 + subjects were given one of three DNA vaccinations for nef, rev or tat, and novel proliferative and CTL responses were generated.

- The nef DNA immunization induced the highest and most consistent CTLp activity, IFNγ production, and IL-6 and IgG responses.
- Highly active antiretroviral treatment (HAART) did not induce new HIV-specific CTL responses but reduced viral load, while DNA vaccination induced new immune responses but did not reduce viral load – thus this is a potentially complementary and promising combination.

**HXB2 Location** Rev **Author Location** Rev

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA HIV component: Nef, Rev, Tat Adjuvant: CpG immunostimulatory sequence (ISS)

Species (MHC) human

Keywords review, Th1

References Calarota & Wahren 2001

This review discusses the cellular immune response, and comments on CpG induction of Th1 cytokines and enhanced immune responses, and HIV-1 DNA vaccine boosting of CTL and Th proliferative responses in asymptomatic HIV+ individuals.

HXB2 Location Rev Author Location Rev Epitope Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade MN HIV component: Env,

Rev Adjuvant: Bupivacaine

Species (MHC) human

**Keywords** early-expressed proteins **References** MacGregor *et al.* 2002

- A phase I clinical trial of a HIV-1 Env and Rev DNA vaccine
  with a CMV promoter was conducted and Th proliferative,
  CTL and Elispot responses monitored. The construct was
  modified for safety and included no LTRs or packaging signals.
  The vaccine strategy was safe, and elicited strong CD4-T cell
  responses, but not CD8 T-cell responses. Rev elicited strong
  Th responses, and is a early produced protein so may confer
  advantages.
- With a 300 ug dose, 4/6 individuals had a lymphocyte proliferation (LP) responses to gp120, 3/6 to Rev.
- With a 1000 ug dose, 4/6 individuals had a LP and 2/6 had IFNgamma Elispot responses to gp160; 3/6 had LP, and 4/6 had IFNgamma Elispot responses to Rev.
- No responses to three specific CTL epitopes were observed by Elispot in individuals with appropriate HLA. Some cytotoxic activity against whole protein was observed that was CD4+ T-cell mediated.

## III-B-13 Vpu Helper/CD4+ T-cell epitopes

HXB2 Location Vpu (19–34) Author Location Vpu (19–34)

**Epitope** AIVVWSIVLIEYRKIL **Immunogen** HIV-1 infection

Species (MHC) human

References Ranki et al. 1997

• T-cell response to this epitope persisted after seroreversion.

HXB2 Location Vpu Author Location Vpu Epitope

Immunogen vaccine

Vector/Type: DNA HIV component: Nef,

Vif, Vpu

**Species (MHC)** mouse (H-2<sup>d</sup>)

**Keywords** subtype comparisons, Th1 **References** Ayyavoo *et al.* 2000

 Splenocytes from BALB/c mice immunized with pVVN-P DNA were incubated with Vif, Vpu or Nef antigens for 3 days and assayed for IL-4 and IFNγ levels.

- Antigen stimulation increased IFNγ production in pVVN-P immunized mice, indicating a Th1 response.
- IL-4 production was not significantly changed after antigen stimulation compared to control levels.
- Cross-clade CTL activity was also observed: A, B clade, CRF01(AE) clade antigens could serve as targets for the B clade immunization stimulated CTL – an HIV-1 AC recombinant, however, did not stimulate a CTL response, but was expressed at lower levels on the target cell.

## III-B-14 gp160 Helper/CD4+ T-cell epitopes

**HXB2 Location** gp160 (30–51)

**Author Location** gp120 (30–51 IIIB)

Epitope ATEKLWVTVYYGVPVWKEATTT?

Epitope name A1 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 2/15 responders recognized this peptide, mean SI = 4.6.

HXB2 Location gp160 (32–44) Author Location gp120 (39–51) Epitope EQLWVTVYYGVPV

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>bxk</sup>)

References Sastry & Arlinghaus 1991

 Peptides induced T-cell proliferative response to immunizing peptide and to gp160. HXB2 Location gp160 (38–48) Author Location gp120 (45–55) Epitope VYYGVPVWKEA Immunogen vaccine

Vector/Type: peptide

**Species** (MHC) mouse (H-2<sup>bxk</sup>, H-2<sup>sxd</sup>) **References** Sastry & Arlinghaus 1991

 Peptides induced T-cell proliferative response to immunizing peptide and to gp160.

HXB2 Location gp160 (38–48) Author Location Env (45–55) Epitope VYYGVPVWKEA Immunogen vaccine

Vector/Type: peptide

Species (MHC) macaque

References Nehete et al. 1993

- Synthetic peptide derived from conserved region of the HIV-1 envelope that stimulates a proliferative response in mice.
- Proliferative response to this peptide was observed in 3/3 immunized rhesus monkeys.

HXB2 Location gp160 (38–48)
Author Location Env (45–55)
Epitope VYYGVPVWKEA
Immunogen HIV-1 infection
Species (MHC) human, chimpanzee
References Nehete et al. 1998b

- Seven out of nine HIV-infected chimpanzees and eight out of seventeen HIV-positive humans exhibited positive proliferative responses to this conserved peptide (peptide 104) – no HIV negative individuals showed a response.
- This peptide, along with 4 other peptides from conserved regions of envelope, can induce proliferative responses to HIV and may be useful for vaccines.
- Peptide 104 elicited proliferative responses in inbred mouse strains and outbred rhesus monkeys in previous study by same group.

HXB2 Location gp160 (41–54) Author Location gp120 (48–61) Epitope GVPVWKEATTLFC Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>sxd</sup>)

References Sastry & Arlinghaus 1991

• Peptides induced T-cell proliferative response to immunizing peptide and to gp160.

HXB2 Location gp160 (41–54) Author Location Env (48–60) Epitope GVPVWKEATTLFC

Immunogen vaccine

Vector/Type: peptide

Species (MHC) macaque

References Nehete et al. 1993

- Synthetic peptide derived from conserved region of the HIV-1 envelope that stimulates a proliferative response in mice.
- Despite the proliferative response to this peptide in mice, no response was observed in 3 rhesus monkeys.

HXB2 Location gp160 (41–60) Author Location gp120 (40–59 89.6) Epitope GVPVWREATTTLFCASDAKA

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse (H-2<sup>d</sup>)

Keywords immunodominance
References Dai *et al.* 2001

- Promiscuous immunodominant epitopes in gp120 were mapped by overlapping peptides in CBA/J H-2<sup>k</sup> and BALB/c H-2<sup>d</sup> mice, and all were found to be in the outer domain, proximal to regions of structural disorder indicated by the crystal structure or by sequence divergence.
- This peptide was recognized by 10/10 BALB/c with an average SI of 6.4, the strongest reaction among BALB/c mice, but not by CBA/J mice, but recognized well not by CBA/J mice, so is considered to be uniquely immunodominant for H-2<sup>d</sup>
- Uniquely immunodominant sequences tended to be in the inner domain of the protein.

HXB2 Location gp160 (41–60) Author Location gp120 (40–59 89.6) Epitope GVPVWREATTTLFCASDAKA

Epitope name Peptide 2

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

**Species (MHC)** mouse **Donor MHC** H-2k, H2-d

**Keywords** epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was highly reactive in 10/10 BALB/c mice tested, but only in 5/10 CBA/J mice.

 $\textbf{HXB2 Location} \ \ gp160 \ (42\text{--}61)$ 

**Author Location** gp120 (42–61 IIIB)

Epitope VPVWKEATTTLFCASDAKAY?

Epitope name A2 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

• Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.

- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 2/15 responders recognized this peptide, mean SI = 6.6.

**HXB2 Location** gp160 (52–71) **Author Location** gp120 (52–71 IIIB)

Epitope LFCASDAKAYDTEVHNVWAT?

Epitope name A3 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 3/15 responders recognized this peptide, mean SI = 4.3.

**HXB2 Location** gp160 (61–80) **Author Location** gp120 (60–79 89.6)

**Epitope** YDTEVHNVWATHACVPTDPN

**Epitope name** Peptide 4 **Immunogen** vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

**Species (MHC)** mouse **Donor MHC** H-2k, H2-d

Keywords epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was highly reactive in 4/10 BALB/c mice tested, but only in 1/10 CBA/J mice.

**HXB2 Location** gp160 (62–80)

Author Location gp120 (62–80 IIIB)

Epitope DTEVHNVWATHACVPTDPN?

Epitope name A4 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 1/15 responders recognized this peptide, SI = 3.5.

**HXB2 Location** gp160 (62–81) **Author Location** gp120 (MN)

Epitope DTEVHNVWATQACVPTDPNP

Epitope name DP20

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DR)

Assay type Cytokine production, proliferation, CD4 Tcell Elispot - IFN $\gamma$ , Intracellular cytokine staining

**Keywords** HAART, ART, acute/early infection, crosspresentation by different HLA

References Malhotra et al. 2003

- 92 acute- or early-HIV infected subjects were tested for Envand Gag-specific Th responses. There was an overall low probability of detecting HIV-1-specific Th responses, Env responses were rare and (5%) and only detected very early in infection, probably reflecting a low frequency of response and impaired lymophoproliferative capacity, not viral escape. Gagspecific Th responses were observed more frequently (16%), were found throughout the acute and early infection phases, and predominated after ARV therapy.
- This epitope is one of six Env-specific Th responses detected in one patient 11 days post-infection that were studied in detail. The CD4+ clones showed MHC-restricted cytotoxicity and secreted high levels of cytokines when stimulated. These Th reponses were not detected at subsequent time points, despite the Env specific Th cells being maintained as detected by TCR clonal transcripts, and the epitope sequence being maintained in Env.
- This peptide showed bound to HLA-DRB1\*0101.

**HXB2 Location** gp160 (65–75)

**Author Location** gp120 (72–82)

Epitope AHKVWATHACV

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>bxk</sup>, H-2<sup>sxd</sup>)

**References** Sastry & Arlinghaus 1991

 Peptides induced T-cell proliferative response to immunizing peptide and to gp160.

**HXB2 Location** gp160 (74–85)

Author Location gp120 (81–92)

**Epitope** CVPTNPVPQEVV

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>bxk</sup>, H-2<sup>sxd</sup>)

References Sastry & Arlinghaus 1991

 Peptides induced T-cell proliferative response to immunizing peptide and to gp160.

HXB2 Location gp160 (74–85) Author Location gp120 (74–85 LAI) Epitope CVPTDPNPQEVV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

HXB2 Location gp160 (80–99)

Author Location gp120 (51–70 HXB2)

Epitope NPQEVVLVNTENFNMWKND
Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human Keywords TCR usage References Li Pira et al. 1998

- Clonal heterogeneity was broad for a recall response to tetanus toxoid or PPD, but oligoclonal to primary HIV antigens, dominated in this case by TCR Vβ 13 usage.
- Donor of PBMC that recognized this epitope had HLA-DR alleles 2 and 7.

**HXB2 Location** gp160 (81–100) **Author Location** gp120 (80–99 89.6)

Epitope PQEVVLGNVTENFNMWKNNM

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

**Species** (MHC) mouse (H-2<sup>k</sup>) **Keywords** immunodominance **References** Dai *et al.* 2001

- Promiscuous immunodominant epitopes in gp120 were mapped by overlapping peptides in CBA/J H-2<sup>k</sup> and BALB/c H-2<sup>d</sup> mice, and all were found to be in the outer domain, proximal to regions of structural disorder indicated by the crystal structure or by sequence divergence.
- This peptide was recognized by 10/10 CBA/J with an average SI of 8.2, and not by BALB/c mice, so is considered to be uniquely immunodominant for H-2<sup>k</sup>
- Uniquely immunodominant sequences tended to be in the inner domain of the protein.

HXB2 Location gp160 (81–100)
Author Location gp120 (80–99 89.6)
Epitope PQEVVLGNVTENFNMWKNNM

**Epitope name** Peptide 6 **Immunogen** vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse Donor MHC H-2k

Keywords epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was not reactive in any BALB/c mice tested (0/10), but was highly reactive in all (10/10) CBA/J mice.

HXB2 Location gp160 (81–101) Author Location gp120 (81–101 IIIB)

Epitope PQEVVLVNVTENFNMWKNDMV?

Epitope name B1 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 2/15 responders recognized this peptide, mean SI = 5.1.

**HXB2 Location** gp160 (92–101)

Author Location gp120 (90-100 W6.ID)

**Epitope** YFNMWKNNMV **Immunogen** vaccine

Vector/Type: protein Strain: B clade W61D HIV component: gp120 Adjuvant: MPL-SE adjuvant, OS21

Species (MHC) human

References Jones et al. 1999

- An HIV seronegative volunteer was vaccinated with rgp120 and a QS21/MPL adjuvant and HIV-1 specific T-cell lines were isolated.
- One T-cell clone reacts with two overlapping peptides, and the region of overlap is: YFNMWKNNMV.
- The first 20-mer peptide that this clone reacts with is PQEVVL-GNVTEYFNMWKNNMV, and the IIIB version of this peptide does not induce proliferation in the T-cell line that responds to the W61D version: IIIB: pqevvlVnvteNfDmwknDmv.

HXB2 Location gp160 (92–111)

Author Location gp120 (92–111 W6.ID)

Epitope YFNMWKNNMVDQMHEDIISL

Immunogen vaccine

Vector/Type: protein Strain: B clade W61D HIV component: gp120 Adjuvant: MPL-SE

adjuvant, QS21

Species (MHC) human

References Jones et al. 1999

- An HIV seronegative volunteer was vaccinated with rgp120 and a QS21/MPL adjuvant and HIV-1 specific T-cell lines were isolated.
- The IIIB version of this peptide does not induce proliferation in the T-cell line that responds to the W61D version of the peptide NfDmwknDmvEqmhediisl.
- Six T-cell lines react with this peptide, but some of these can also be stimulated by other gp120 peptides located in different regions of gp120.

**HXB2 Location** gp160 (101–126) **Author Location** gp120 (101–126)

Epitope VEQMHEDIISLWDQSLKPCVKLTPLC

Immunogen vaccine

Vector/Type: protein HIV component: gp160

**Species (MHC)** mouse (H-2<sup>k</sup>) **References** Sjolander *et al.* 1996

• Study showing that T-cell determinants from glycoproteins can be dependent on the glycosylation of the protein.

HXB2 Location gp160 (102–114) Author Location gp120 (109–121) Epitope EQMHEDIISLWDQ

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>bxk</sup>)

References Sastry & Arlinghaus 1991

 Peptides induced T-cell proliferative response to immunizing peptide and to gp160.

HXB2 Location gp160 (102–116) Author Location gp160 (109–123 IIIB) Epitope EQMHEDIISLWDQSL

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

**References** Berzofsky et al. 1991b; Berzofsky et al. 1991a

- B10.D2 (H-2A<sup>d</sup>, E<sup>d</sup>) and B10.A(R5) (H-2A<sup>b</sup>, E<sup>b</sup>) mice immunized with rec gp160 showed a proliferative response to EQMHEDIISLWDQSL.
- EQMHEDIISLWDQSLKPCVK encompasses several murine Th epitopes including HEDIISLWDQSLK and is referred to as a "multideterminant region" or cluster peptide.

HXB2 Location gp160 (102–116) Author Location gp120 (109–123 IIIB) Epitope EQMHEDIISLWDQSL

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160

Species (MHC) mouse (H-2<sup>d</sup>, H-2<sup>i5</sup>)
References Hale *et al.* 1989

 Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

**HXB2 Location** gp160 (102–121) **Author Location** gp160 (109–128 IIIB)

Epitope EQMHEDIISLWDQSLKPCVK

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** human, mouse (H-2<sup>k</sup>, H-2<sup>s</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- EQMHEDIISLWDQSLKPCVK encompasses several murine
  Th epitopes and is referred to as a "multideterminant region"
  or cluster peptide.
- Six multideterminant region cluster peptides were evaluated Th responses in different MHC/HLA backgrounds after vaccination of mice with gp160, or in infected people.
- This cluster peptide elicited proliferative responses in cells from vaccinated B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>) and B10.S(9R) mice (H-2A<sup>s</sup>, E<sup>s</sup>), while shorter peptides from within this region stimulated H-2<sup>k</sup>, H-2<sup>d</sup> and H-2<sup>b</sup> responses, but not H-2<sup>s</sup>
- IL-2 production was observed in response to this peptide in 64% (23/36) of asymptomatic HIV-infected individuals.

HXB2 Location gp160 (102–121)

Author Location gp120 (102–121 IIIB)

Epitope EQMHEDIISLWDQSLKPCVK?

Epitope name B3 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 1/15 responders recognized this peptide, SI = 5.9.

**HXB2 Location** gp160 (105–117) **Author Location** gp120 (112–124 IIIB)

Epitope HEDIISLWDQSLK

Epitope name T2

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>k</sup>) **References** Hale *et al.* 1989

• Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

HXB2 Location gp160 (105–117) Author Location gp160 (112–124 IIIB) Epitope HEDIISLWDQSLK

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- B10.BR (H-2A<sup>k</sup>, E<sup>k</sup>) mice immunized with rec gp160 showed a strong proliferative response to three overlapping peptides, QMHEDIISLWDQSL, HEDIISLWDQSLK, and DIISLWDQS-LKPCVK, and HEDIISLWDQSLK is common to between them.
- EQMHEDIISLWDQSLKPCVK encompasses several murine Th epitopes including HEDIISLWDQSLK and is referred to as a "multideterminant region" or cluster peptide.

HXB2 Location gp160 (105–117) Author Location gp120 (112–124 BH10) Epitope HEDIISLWDQSLK

Epitope name T2

**Immunogen** computer prediction **Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>s</sup>) **References** Cease *et al.* 1987

 1 of 2 functional epitopes identified using an amphipathic helix epitope prediction algorithm.

HXB2 Location gp160 (105–117) Author Location gp120 (112–124 IIIB) Epitope HEDIISLWDQSLK

Epitope name T2

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1997

• Used in a study of pentoxifylline's influence on HIV specific T-cells.

HXB2 Location gp160 (105–117) Author Location gp120 (112–124 BH10) Epitope HEDIISLWDQSLK

Epitope name T2 Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: gp160

Species (MHC) human

References Berzofsky et al. 1988

 Proliferative response to T1 and T2 peptides in 14 immunized, uninfected humans.

HXB2 Location gp160 (105–117) Author Location gp120 (112–124 IIIB) Epitope HEDIISLWDQSLK

Epitope name T2

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1989

IL-2 production detection of Th lymphocytes from asymptomatic HIV-positive individuals.

HXB2 Location gp160 (105–117) Author Location gp120 (112–124 IIIB) Epitope HEDIISLWDQSLK

Epitope name T2

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1991a

 Peptides stimulate Th cell function and CTL activity in similar patient populations.

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HXB2 Location gp160 (105–117)
Author Location gp120 (112–124)
Epitope HEDIISLWDQSLK
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Epitope name T2 Immunogen vaccine

*Vector/Type:* protein *Strain:* B clade IIIB *HIV component:* gp160

Species (MHC) human

References Clerici et al. 1991b

• Immunizing uninfected individuals with rgp160 results in stronger Th response than does natural infection.

HXB2 Location gp160 (105–117) Author Location gp120 (112–124 IIIB) Epitope HEDIISLWDQSLK Epitope name T2

Immunogen

Species (MHC) human

References Clerici et al. 1992

• Cell-mediated immune response to HIV-1 peptides in HIV-1

exposed seronegative men.

HXB2 Location gp160 (105–117)

Author Location gp120 (112–124 IIIB)

Epitope HEDIISLWDQSLK

Epitope name T2 Immunogen vaccine

Vector/Type: peptide prime with protein boost Strain: B clade IIIB HIV component: gp160

Species (MHC) macaque

References Hosmalin et al. 1991

Peptide priming to induce T-cell help enhances antibody response to gp160 immunization.

HXB2 Location gp160 (105–117) Author Location gp120 (112–124 IIIB) Epitope HEDIISLWDQSLK

Epitope name T2 Immunogen

Species (MHC) human

**References** Pinto *et al.* 1995

• CTL activity analyzed in parallel with Th reactivity in exposed

but uninfected health care workers.

HXB2 Location gp160 (105–117) Author Location gp120 (112–124 IIIB) Epitope HEDIISLWDQSLK

Epitope name T2

Immunogen HIV-1 infection

Species (MHC) human

References Kaul et al. 1999

- Kenyan sex workers that remained seronegative were found to frequently have HIV-env peptide specific Th responses detected by an IL-2 assay (11/20 cases) and mucosal genital tract anti-HIV IgA (16/21 cases)
- Helper epitopes used in this study were noted to be previously described Clerici et al. [1989], and were not explicitly described in Kaul et al. [1999]

**HXB2 Location** gp160 (105–117)

**Author Location** gp120

Epitope HEDIISLWDQSLK

Epitope name T2

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human

**Keywords** subtype comparisons, responses in children,

mother-to-infant transmission

References Kuhn et al. 2001a

- In a S. African perinatal transmission study, 33% (33/86) of cord blood samples from infants with seropositive mothers produced T-helper responses (measured by a bioassay measuring IL2 production in a murine cell line and confirmed with a proliferation assay) against a peptide cocktail containing Th epitopes P18 MN, P18 IIIB, T1, T2, and TH4.
- The mothers were predominantly infected with subtype C, but the T help response was detectable in a number of cord blood samples despite using peptides based on B subtype reagents.
- 3/33 infants with cord blood T help responses to Env were infected in utero, 2/33 were lost to follow up, and 28/33 were not infected. 6/53 of the infants with cord blood that was unresponsive to Env peptide stimulation were infected before delivery, and 8/47 contracted HIV intrapartum or via breastfeeding.
- Measurable HIV specific T help responses elicited in the immunologically immature newborn, possibly in response to in utero exposure, are associated with a protective natural immunity that helps block mother-infant transmission of HIV-1.

**HXB2 Location** gp160 (105–117) **Author Location** Env (112–124 IIIB)

Epitope HEDIISLWDQSLK

Epitope name T2

Subtype B

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC)

Assay type Cytokine production

Keywords mother-to-infant transmission

References Clerici et al. 1993a

- Cord blood samples in 8/23 infants with HIV+ mothers showed IL-2 production in response to peptides from HIV-1 gp1260, demonstrating that Th cell priming to HIV env determinants can occur in utero. HIV-specific Th immunity in utero may be protective, as none of the 8 with HIV-1 specific Th activity became infected, while 3/15 infants with no detectable Th activited were infected.
- PBL from 10/21 of the mothers showed HIV-1 specific Th activity through IL-2 production in response to Env peptides.

**HXB2 Location** gp160 (105–117)

**Author Location** Env (IIIB)

Epitope HEDIISLWDQSLK

**Epitope name** T2

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC)

**Assay type** Cytokine production **References** Clerici *et al.* 1994a

- Six of eight HIV-exposed health care workers had transient HIV-specific T-helper responses after percutaneous exposure to HIV, responding to two or more Env peptides among the set T1, T2, Th4.1, P18IIIB and P18MN. Responses were lost after 12-56 weeks. The HIV-specific Th responses occurred without seroconversion or PCR evidence for infection.
- Six of the eight HIV-exposed individuals responded to two or more peptides, whereas, only one individual exposed to HIV-negative blood responded to one peptide.

**HXB2 Location** gp160 (105–117)

**Author Location** HIV-1 (IIIB)

Epitope HEDIISLWDQSLK

Epitope name T2

Subtype B

Immunogen HIV-1 infection

Species (MHC)

Assay type Cytokine production

References Clerici et al. 1994b

IL-10-specific mRNA was upregulated in PBMC from asymptomatic, HIV-infected (HIV+) patients, and was particularly high in those with severely compromised Th cells function. Th response to HIV peptides *in vitro* could be restored by IL-10 Ab.

**HXB2 Location** gp160 (105–117)

Author Location Env (112-124)

Epitope HEDIISLWDQSLK

Epitope name T2

Immunogen HIV-1 infection

Species (MHC) human

Assay type proliferation

Keywords responses in children, mother-to-infant trans-

mission

References Kuhn et al. 2001b

- Thelper proliferative responses in cord blood at delivery to a cocktail of HIV envelope peptides were infrequent (1/41) among infants whose HIV-positive mothers received antiretroviral prophylaxis at delivery to prevent transmission. Responses were 10 times more frequent among infants of HIV-seropositive women who had no antiretroviral treatment (7/29). Reductions of HIV-specific responses occurred despite persistance of detectable HIV RNA in the mothers at delivery.
- The reduction of Th responses in newborns raises the possibility that anti-retrovial exposure during pregnancy may block subsequent immune protection. The authors point two relevant citations that showed ARV exposed infants had a more rapid course of progression (Kuhn *et al.*, JID 182:104 (2000)), but were at no greater risk of infection due to subsequent breast feeding (Ditrame *et al.*, Lancet 354:2050 (1999)).

 $\textbf{HXB2 Location} \hspace{0.1cm} gp160 \hspace{0.1cm} (105\text{--}117)$ 

Author Location Env (gp160) (105–117)

Epitope HEDIISLWDQSLK

Epitope name TH2

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Country South Africa.

Assay type proliferation

recognition or cross-neutralization

References Meddows-Taylor et al. 2004

- Viral isolates (gp160) from 16 vertically HIV-1 infected children (8 T-helper cell Env responders and 8 non-responders) were analyzed for variation after comparison with Env peptide sequences (conserved regions T1, T2 and TH4.1, hypervariable regions P18 MN and P181 IIIB) used for in vitro stimulation.
- T2 was the most conserved of the 5 peptides studied.
- No correlation between the age, clinical category (mild or severe), HIV-1 viral load and the degree of epitope variation was established.

HXB2 Location gp160 (105-123) **Author Location** gp120 (112–130 IIIB) Epitope HEDIISLWDQSLKPCVKLT

Immunogen

Species (MHC) human

References Furci et al. 1997

• 9/11 exposed-uninfected individuals in this study had a proliferative response to a C5 peptide, but none reacted with this previously defined epitope.

HXB2 Location gp160 (108-119) Author Location gp120 (108–119 LAI) Epitope IISLWDQSLKPC

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

**HXB2 Location** gp160 (110–125) Author Location gp120 (110-125) Epitope SLWDQSLKPCVKLTPL Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression References Caruso et al. 1997

- · As HIV-1-infected individuals progress to disease, T-cells show reduced ability to proliferate in response to HIV antigen, but retain the ability to express the activation antigens CD25 and CD71.
- The ability to express activation markers in response to HIV is retained, but the response to tetanus toxoid recall antigen is lost.
- This study investigated CD25 and CD71 expression in PBMC from patients at various stages of progression, measuring the response to in vitro stimulation by peptide cocktail containing four antigenic Env peptides, or p17 and p24.

**HXB2 Location** gp160 (111–123) Author Location gp120 (118–130) Epitope LWDQSLKPCVKLT

Immunogen vaccine

Vector/Type: peptide

Species (MHC) macaque

References Nehete et al. 1993

• Synthetic peptide derived from conserved region of the HIV-1 envelope that stimulates a proliferative response in mice.

**Keywords** responses in children, variant cross- • Proliferative response to this peptide was observed in 3/3 immunized rhesus monkeys.

HXB2 Location gp160 (112-130)

Author Location gp120 (112-130 IIIB)

Epitope WDQSLKPCVKLTPLCVSLK?

Epitope name B4 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- · After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 3/15 responders recognized this peptide, average SI = 4.4.

**HXB2 Location** gp160 (112–141) **Author Location** gp120 (112–141 NL43)

Epitope WDQSLKPCVKLTPLCVSLKCTDLGNATNTN

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: gp120, gp160

Species (MHC) human

References Sitz et al. 1999

- There was a great breadth of proliferative response to env peptides in 19 HIV-1 infected rgp160 and 17 HIV-1 infected rgp120 vaccine recipients.
- Over 35% of vaccinees had a stimulation index of greater than 5 to this peptide.

HXB2 Location gp160 (115-126)

Author Location gp120 (115-126 LAI)

**Epitope** SLKPCVKLTPLC

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

**HXB2 Location** gp160 (115–129)

Author Location gp120 (115–129 LAI)

**Epitope** SLKPCVKLTPLCVSL

Subtype B

Immunogen peptide-HLA interaction

Species (MHC) human (HLA-DR)

Keywords binding affinity

References Gaudebout et al. 1997

- Peptide bound to both HLA-DR\*1101 and HLA-DR\*0401 with high affinity.
- Because of the distinctive binding pockets of HLA-DR\*1101 and HLA-DR\*0401, peptides that bound both were considered candidates for promiscuous HLA-DR binding.

HXB2 Location gp160 (121-140) **Author Location** gp120 (120–139 89.6) Epitope KLTPLCVTLNCTNLNITKNT

Epitope name Peptide 10 Immunogen vaccine

> Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse Donor MHC H-2d

Keywords epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was highly reactive in 5/10 BALB/c mice tested, but not in and (0/10) CBA/J mice.

HXB2 Location gp160 (121-141)

Author Location gp120 (131–151 IIIB)

**Epitope** KLTPLCVSLKCTDLKNDTNTN?

Epitope name C1 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- · After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 3/15 responders recognized this peptide, average SI = 3.9.

HXB2 Location gp160 (122–141)

Author Location gp120 (121–140 MN)

**Epitope** LTPLCVTLNCTDLRNTTNTN

Epitope name 1931

Subtype B

Immunogen vaccine

Vector/Type: DNA, protein Strain: B clade MN HIV component: gp120 Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) guinea pig

Keywords vaccine-specific epitope characteristics, Th1

References Chattergoon et al. 2002

• Hartley guinea pigs were intradermally injected with either recombinant protein or plasmid expressed gp120 and monitored for delayed type hypersensivity (DTH) responses after vaccination, which are related to Th1 T-cell responses. CFA did not augment responses in animals vaccinated with plasmid.

- A total of 7 gp120 peptides elicited a delayed type hypersensivity (DTH) response after vaccination, out of a set of 60 overlapping peptides that spanned gp120. The vaccine delivery system, DNA versus rec protein, resulted in the recognition of distinct peptides.
- 3/5 animals vaccinated with rec gp120 responded by DTH to this peptide, while 0/6 vaccinated with plasmid gp120 DNA responded.

HXB2 Location gp160 (122-141)

Author Location gp120 (122-141 IIIB)

**Epitope** LTPLCVSLKCTDLKNDTNTN?

Epitope name B5

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- 1/15 responders recognized this peptide, SI = 3.1.

HXB2 Location gp160 (136–155)

Author Location gp120 (141–160 MN)

Epitope NSTAWNNSNSEGTIKGGEMK

Epitope name 1932

Subtype B

Immunogen vaccine

Vector/Type: DNA, protein Strain: B clade MN HIV component: gp120 Adjuvant:

Complete Freund's Adjuvant (CFA)

Species (MHC) guinea pig

**Keywords** vaccine-specific epitope characteristics, Th1

References Chattergoon et al. 2002

- Hartley guinea pigs were intradermally injected with either recombinant protein or plasmid expressed gp120 and monitored for delayed type hypersensivity (DTH) responses after vaccination, which are related to Th1 T-cell responses. CFA did not augment responses in animals vaccinated with plasmid.
- A total of 7 gp120 peptides elicited a delayed type hypersensivity (DTH) response after vaccination, out of a set of 60 overlapping peptides that spanned gp120. The vaccine delivery system, DNA versus rec protein, resulted in the recognition of distinct peptides.
- 4/5 animals vaccinated with rec gp120 responded by DTH to this peptide, as did 2/6 vaccinated with plasmid gp120 DNA.

HXB2 Location gp160 (138-159)

Author Location gp120 (141–160 W6.ID)

Epitope TTSNGWTGEIRKGEIKNCSF

Immunogen vaccine

Vector/Type: protein Strain: B clade W61D HIV component: gp120 Adjuvant: MPL-SE

adjuvant, QS21

Species (MHC) human

References Jones et al. 1999

- An HIV seronegative volunteer was vaccinated with rgp120 and a QS21/MPL adjuvant and HIV-1 specific T-cell lines were isolated.
- The IIIB version of this peptide does not induce proliferation in the T-cell line that responds to the W61D version of the peptide: IIIB: ttsnSSGRMIMEgeikncsf.

**HXB2 Location** gp160 (142–161) **Author Location** gp120 (142–161 IIIB)

Epitope SSSGRMIMEKGEIKNCSFNI?

Epitope name C2

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** immunodominance **References** Geretti *et al.* 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- Five peptides were recognized most frequently: C2 (aa 142-161), C3 (aa 152-171), C5 (aa 172-191), E5 (aa 272-291) and G4 (aa 380-393). The first three were in or near V2, the other two were proximal to the V3 and V4 loops.
- 4/15 responders recognized this immunodominant peptide, average SI = 4.3.

**HXB2 Location** gp160 (147–168)

**Author Location** gp120 (152–173 NL43)

Epitope MMMEKGEIKNCSFNISTSIRGK

Immunogen vaccine

*Vector/Type:* protein *Strain:* B clade NL43 *HIV component:* gp120, gp160

Species (MHC) human

References Sitz et al. 1999

- There was a great breadth of proliferative response to env peptides in 19 HIV-1 infected rgp160 and 17 HIV-1 infected rgp120 vaccine recipients.
- Over 50% of vaccinees had a stimulation index of greater than 5 to this peptide.

**HXB2 Location** gp160 (152–171)

Author Location gp120 (152–171 IIIB)

Epitope GEIKNCSFNISTSIRGKVQK?

Epitope name C3

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** immunodominance **References** Geretti *et al.* 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- Five peptides were recognized most frequently: C2 (aa 142-161), C3 (aa 152-171), C5 (aa 172-191), E5 (aa 272-291) and G4 (aa 380-393). The first three were in or near V2, the other two were proximal to the V3 and V4 loops.
- 4/15 responders recognized this immunodominant peptide, average SI = 4.4.

HXB2 Location gp160 (155–169)

Author Location Env (UG92005)

**Epitope** KNCSFNITTELIDKK

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the V2 region of UG92005 (UG, clade D) and the hybridoma that recognized it used V $\beta$ 5.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the
  first 38 amino acids contributed by BH10 and the rest of gp120
  and gp41 by the vaccine strain, the DNA construct is in the
  pJW4303 vector with a CMV promotor, and the purified protein
  is expressed from the pJW4303 vector transfected into CHO-K1
  cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (155-169) Author Location gp120 (160–174 LAI)

**Epitope** KNCSFNISTSIRGKV

Subtype B Immunogen

**Species (MHC)** human (HLA-DR) **Keywords** binding affinity

References Gaudebout et al. 1997

- Peptide binds to both HLA-DR\*1101 and HLA-DR\*0401 with IL-2 production was a more sensitive and well-preserved meahigh affinity.
- Because of the distinctive binding pockets of HLA-DR\*1101 and HLA-DR\*0401, peptides that bound both were considered candidates for promiscuous HLA-DR binding.

HXB2 Location gp160 (159-178) **Author Location** gp120 (160–179 89.6) Epitope FYITTSIRNKVKKEYALFNR

Epitope name Peptide 14 Immunogen vaccine

> Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse Donor MHC H-2k, H2-d

Keywords epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was highly reactive in 6/10 BALB/c mice tested, and in 4/10 CBA/J mice.

HXB2 Location gp160 (162-181) Author Location gp120 (162–181 IIIB)

Epitope STSIRGKVQKEYAFFYKLDI

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB Author Location gp120 (141–160 W6.ID)

HIV component: Env

Species (MHC) macaque

References Lekutis et al. 1997

• HIV-1 env DNA vaccine induced Th cell response to this epitope in a rhesus monkeys.

**HXB2 Location** gp160 (162–182)

Author Location gp120 (162–182 IIIB)

Epitope STSIRGKVQKEYAFFYKLDII?

Epitope name C4 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- sure of Th function than proliferation.
- 1/15 responders recognized this peptide, SI = 3.3.

HXB2 Location gp160 (166-185) Author Location gp120 (MN)

Epitope RDKMQKEYALLYKLDIVSID

Epitope name RD20 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type Cytokine production, proliferation, CD4 Tcell Elispot - IFNγ, Intracellular cytokine

staining

Keywords HAART, ART, acute/early infection

References Malhotra et al. 2003

- 92 acute- or early-HIV infected subjects were tested for Envand Gag-specific Th responses. There was an overall low probability of detecting HIV-1-specific Th responses, Env responses were rare and (5%) and only detected very early in infection, probably reflecting a low frequency of response and impaired lymophoproliferative capacity, not viral escape. Gagspecific Th responses were observed more frequently (16%), were found throughout the acute and early infection phases, and predominated after ARV therapy.
- This epitope is one of six Env-specific Th responses detected in one patient 11 days post-infection that were studied in detail. These Th reponses were not detected at subsequent time points, despite the Env specific Th cells being maintained as detected by TCR clonal transcripts, and the epitope sequence being maintained in Env.

HXB2 Location gp160 (169–189)

Epitope VQKEYALFYNLDVVPIDDDNA

Immunogen vaccine

Vector/Type: protein Strain: B clade W61D HIV component: gp120 Adjuvant: MPL-SE adjuvant, QS21

Species (MHC) human

References Jones et al. 1999

- An HIV seronegative volunteer was vaccinated with rgp120 and a QS21/MPL adjuvant and HIV-1 specific T-cell lines were isolated.
- The IIIB version of this peptide does not induce proliferation in the T-cell line that responds to the W61D version of the peptide -F-K-II-N-TT vqkeyaFfyKldIIpidNdTT.
- · Two T-cell lines react specifically with this peptide.

HXB2 Location gp160 (172–191) Author Location gp120 (172-191 IIIB) Epitope EYAFFYKLDIIPIDNDTTSY Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB

HIV component: Env

Species (MHC) macaque

References Lekutis et al. 1997

 HIV-1 env DNA vaccine induced Th cell response to this epitope in a rhesus monkey.

**HXB2 Location** gp160 (172–191)

Author Location gp120 (172–191 IIIB)

Epitope EYAFFYKLDIIPIDNDTTSY?

Epitope name C5

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** immunodominance **References** Geretti *et al.* 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- Five peptides were recognized most frequently: C2 (aa 142-161), C3 (aa 152-171), C5 (aa 172-191), E5 (aa 272-291) and G4 (aa 380-393). The first three were in or near V2, the other two were proximal to the V3 and V4 loops.
- 4/15 responders recognized this immunodominant peptide, average SI = 7.4.

**HXB2 Location** gp160 (175–189) **Author Location** Env (UG92005)

Epitope LFYKLDVVQIDNSTN

Epitope LFIKLDVVQIDN

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the V2 region of UG92005 (UG, clade D) and the  $V\beta$  usage of the TCR was not determined.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.

- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IAb restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (186–208)

**Author Location** Env

Epitope NDNTSYRLISCNTSVITQACPKV

Epitope name HIV\_env\_DRB0101\_3

Subtype M

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type T-cell Elispot

Keywords computational epitope prediction

References De Groot et al. 2004

- Bioinformatic tools (EpiMatrix and Conservatrix) were employed to predict 9 highly conserved and promiscuous class II restricted T helper epitopes from HIV-1 env sequences for a cross-clade polyepitope vaccine. 10/34 HIV-1 infected patients induced Th1 immune responses to 8/9 study peptides.
- 5/34 subjects tested responded to this peptide.
- Class II peptides containing putative epitopes were selected to favor DRB0101 presentation, as there is a transgenic mouse model for this allele for future vaccine studies. The highest scoring DRB0101 sequence of this peptide was YRLISCNTS.

**HXB2 Location** gp160 (186–215)

**Author Location** gp120 (191–220 NL43)

Epitope NDTTSYTLTSCNTSVITQACPKVSFEPIPI

Immunogen vaccine

*Vector/Type:* protein *Strain:* B clade NL43 *HIV component:* gp120, gp160

Species (MHC) human

References Sitz et al. 1999

- There was a great breadth of proliferative response to env peptides in 19 HIV-1 infected rgp160 and 17 HIV-1 infected rgp120 vaccine recipients.
- Over 30% of vaccinees had a stimulation index of greater than 5 to this peptide.

**HXB2 Location** gp160 (188–207)

**Author Location** gp120 (190–209 89.6)

Epitope NTKYRLISCNTSVITQACPK

Immunogen vaccine

HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

**Species (MHC)** mouse (H-2<sup>k</sup>) Keywords immunodominance References Dai et al. 2001

- Promiscuous immunodominant epitopes in gp120 were mapped by overlapping peptides in CBA/J H-2<sup>k</sup> and BALB/c H-2<sup>d</sup> mice, and all were found to be in the outer domain, proximal to regions of structural disorder indicated by the crystal structure or by sequence divergence.
- This peptide was recognized by 9/10 CBA/J with an average SI of 9.8, one of the two immunodominant peptides in CBA/J mice, and not by BALB/c mice, so is considered to be uniquely immunodominant for H-2k
- Uniquely immunodominant sequences tended to be in the inner domain of the protein.

HXB2 Location gp160 (188-207) Author Location gp120 (89.6)

Epitope NTKYRLISCNTSVITQACPK

Epitope name Peptide 17 Immunogen vaccine

> Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse Donor MHC H-2k, H2-d

Keywords epitope processing, immunodominance

References Dai et al. 2001

• Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.

• This peptide was reactive in only 1/10 BALB/c mice tested, but was one of the most reactive in CBA/J mice, reacting with 9/10 mice.

HXB2 Location gp160 (190-212) **Author Location** Env (185–215)

Epitope SYRLISCNTSVITQACPKVSFEP

Epitope name HIV\_env\_DRB0101\_62

Subtype M

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type T-cell Elispot

**Keywords** computational epitope prediction

References De Groot et al. 2004

- · Bioinformatic tools (EpiMatrix and Conservatrix) were employed to predict 9 highly conserved and promiscuous class II restricted T helper epitopes from HIV-1 env sequences for a cross-clade polyepitope vaccine. 10/34 HIV-1 infected patients induced Th1 immune responses to 8/9 study peptides.
- 1/34 subjects tested responded to this peptide.

Vector/Type: protein Strain: B clade 89.6 • Class II peptides containing putative epitopes were selected to favor DRB0101 presentation, as there is a transgenic mouse model for this allele for future vaccine studies. The highest scoring DR B0101 sequence of this peptide was NTSVITQA.

HXB2 Location gp160 (192–211)

Author Location gp120 (192-211 IIIB)

Epitope KLTSCNTSVITQACPKVSFE?

Epitope name D2

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide
- · After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 1/15 responders recognized this peptide, SI = 3.6.

HXB2 Location gp160 (193–218)

**Author Location** gp120 (193–218)

Epitope LTSCNSVITQACPKVSFEPIPIHYC

Immunogen vaccine

*Vector/Type:* protein HIV component:

gp160

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

References Siolander et al. 1996

• Study showing that T-cell determinants from glycoproteins can be dependent on the glycosylation of the protein.

HXB2 Location gp160 (198-212)

**Author Location** Env (1007)

Epitope TSVITQACPKVSFEP

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete

Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

Keywords subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the C2 region of 1007 (US, clade B) and the V $\beta$  usage of the TCRs for two clonotypes was V $\beta$ 3 and VB8.1-2.
- · C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein

is expressed from the pJW4303 vector transfected into CHO-K1 cells.

- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IAb transfected L cells as targets and  $V\beta$  usage was determined.
- · Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IAb restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (198-215) Author Location Env (1007)

Epitope TSVITQACPKVSFEPIPI

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

Keywords subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the C2 region of 1007 (US, clade B) and the  $V\beta$  usage of the TCR was  $V\beta$ 6.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and  $V\beta$  usage was determined.
- · Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site

proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (199-211)

Author Location gp120 (204–216)

**Epitope** SVITQACSKVSFE

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>bxk</sup>, H-2<sup>sxd</sup>)

References Sastry & Arlinghaus 1991

· Peptides induced T-cell proliferative response in mice representing four haplotypes.

HXB2 Location gp160 (199-211)

**Author Location** Env (204–216)

**Epitope** SVITQACSKVSFE

Immunogen vaccine

Vector/Type: peptide

Species (MHC) macaque

References Nehete et al. 1993

- Synthetic peptide derived from conserved region of the HIV-1 envelope that stimulates a proliferative response in mice.
- A weak or transient proliferative response to this peptide was observed in 3/3 immunized rhesus monkeys.

HXB2 Location gp160 (199–211)

Author Location Env (204-216)

**Epitope** SVITQACSKVSFE

Immunogen HIV-1 infection

Species (MHC) human, chimpanzee

References Nehete et al. 1998b

· HIV-infected chimpanzees and HIV-positive patients show positive proliferative responses to multiple peptides from five conserved regions of the HIV-1 Env.

HXB2 Location gp160 (200-214)

Author Location gp120 (205-219 LAI)

Epitope VITQACPKVSFEPIP

Subtype B

Immunogen peptide-HLA interaction

Species (MHC) human (HLA-DR)

Keywords binding affinity

References Gaudebout et al. 1997

- Peptide binds to both HLA-DR\*1101 and HLA-DR\*0401 with high affinity.
- Because of the distinctive binding pockets of HLA-DR\*1101 and HLA-DR\*0401, peptides that bound both were considered candidates for promiscuous HLA-DR binding.

HXB2 Location gp160 (201–212)

Author Location Env (1007)

Epitope ITQACPKVSFEP

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete

Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

Keywords subtype comparisons, epitope processing, TCR usage

## References Surman et al. 2001

- This epitope is located in the C2 region of 1007 (US, clade B) and the Vβ usage of the TCR was Vβ3.
- The epitope described here is the region of overlap of two 15 mers that were both able to stimulate IL-2 production from the hybridoma (TSVITQACPKVSFEP and ITQACPKVSFEPIPI)
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and  $V\beta$  usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (206–220) **Author Location** Env (1007)

Epitope PKVSFEPIPIHYCAP

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing **References** Surman *et al.* 2001

- This epitope is located in the C2 region of 1007 (US, clade B) and 12 hybridomas recognized the peptide with Vβ usage of Vβ4,6,7,8.1-2,8.3,11,12 and others not determined.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.

- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (206–220)

**Author Location** Env (gp160)

Epitope PKVSFEPIPIHYCAP

Subtype B, D

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: Env Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (H2b)

Assay type Cytokine production, CD4 T-cell Elispot -

IFN $\gamma$ 

**Keywords** vaccine-induced epitopes, variant crossrecognition or cross-neutralization, vaccine

recognition or cross-neutralization, vaccine antigen design

References Zhan et al. 2004

- To develop a polyvalent Env vaccine, subtype specific B and D T-helper epitopes were identified, and mixtures of strain 1007, clade B, or UG92005, clade D envelopes were given to C57BL/6J mice. Mice were intramuscularly immunized with recombinant DNA, then intraperitoneally with rVV and finally with env protein in CFA. A dilution of 1/100 could still elicit type specific responses, and responses persisted with different prime boost combinations.
- Priming with 1007 and UG92005 env's induced both Envspecific (SNNTVGNPIILPCRI and AMYAPPIAGLIQCSS) and cross-reactive (PKVSFEPIPIHYCAP, C2, GNIRQAHCN-VSKAKW, V3-C3) Th responses in murine spleen cells.

**HXB2 Location** gp160 (206–225)

Author Location gp120 (211–230 MN)

Epitope PKISFEPIPIHYCAPAGFAI

Epitope name 1957 Subtype B

Immunogen vaccine

Vector/Type: DNA, protein Strain: B clade MN HIV component: gp120 Adjuvant:

Complete Freund's Adjuvant (CFA)

Species (MHC) guinea pig

**Keywords** vaccine-specific epitope characteristics, Th1

References Chattergoon et al. 2002

- Hartley guinea pigs were intradermally injected with either recombinant protein or plasmid expressed gp120 and monitored for delayed type hypersensivity (DTH) responses after vaccination, which are related to Th1 T-cell responses. CFA did not augment responses in animals vaccinated with plasmid.
- A total of 7 gp120 peptides elicited a delayed type hypersensivity (DTH) response after vaccination, out of a set of 60 overlapping peptides that spanned gp120. The vaccine delivery system, DNA versus rec protein, resulted in the recognition of distinct peptides.
- 5/5 animals vaccinated with rec gp120 responded by DTH to this peptide, as did 2/6 vaccinated with plasmid gp120 DNA.

**HXB2 Location** gp160 (206–230) **Author Location** gp120 (206–230)

Epitope PKVSFEPIPIHYCAPAGFAILKCNN

Immunogen vaccine

Vector/Type: protein HIV component:

gp160

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

References Sjolander et al. 1996

• Study showing that T-cell determinants from glycoproteins can be dependent on the glycosylation of the protein.

HXB2 Location gp160 (208–218) Author Location Env (UG92005) Epitope ITFEPIPIHYC

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing

References Surman et al. 2001

- This epitope is located in the C2 region of UG92005 (UG, clade D) and its was recognized by two hybridomas with  $V\beta$  usage  $V\beta$ 12 and not determined.
- The epitope described here is the region of overlap of two 15 mers that were both able to stimulate IL-2 production from the hybridoma (PKITFEPIPIHYCAP and ITFEPIPIHYCAPAG)
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.

- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (208–222) Author Location Env (UG92005)

**Epitope** ITFEPIPIHYCAPAG **Immunogen** vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the C2 region of UG92005 (UG, clade D) and it was recognized by five hybridomas with  $V\beta$  usage  $V\beta$ 5, 8.2, 12 and not determined.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (208–227) Author Location gp120 (210–229 89.6) Epitope VSFQPIPIHYCVPAGFAMLK

**Epitope name** Peptide 19 **Immunogen** vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

**Species (MHC)** mouse **Donor MHC** H-2k, H2-d

**Keywords** epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was reactive in 6/10 BALB/c mice tested, and in 6/10 CBA/J mice.

HXB2 Location gp160 (209–220)

Author Location gp120 (MN)

Epitope SFEPIPIHYCAP

Epitope name SP12

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DR)

Assay type Cytokine production, proliferation, CD4 T-

cell Elispot - IFN  $\gamma$ , Intracellular cytokine

staining

**Keywords** HAART, ART, vaccine-specific epitope characteristics, acute/early infection, cross-

presentation by different HLA

References Malhotra et al. 2003

- 92 acute- or early-HIV infected subjects were tested for Envand Gag-specific Th responses. There was an overall low probability of detecting HIV-1-specific Th responses, Env responses were rare and (5%) and only detected very early in infection, probably reflecting a low frequency of response and impaired lymophoproliferative capacity, not viral escape. Gag-specific Th responses were observed more frequently (16%), were found throughout the acute and early infection phases, and predominated after ARV therapy.
- This epitope is one of six Env-specific Th responses detected in one patient 11 days post-infection that were studied in detail. The CD4+ clones showed MHC-restricted cytotoxicity and secreted high levels of cytokines when stimulated. These Th reponses were not detected at subsequent time points, despite the Env specific Th cells being maintained as detected by TCR clonal transcripts, and the epitope sequence being maintained in Env
- Seven out of 12 clones recognized this conserved C3 region of gp120. Clone one was mapped to the optimal epitope and was found to be presented by HLA-DR. The peptide showed promiscuous binding to DRB1\*0101, DRB1\*0401, DRB1\*1302, DRB1\*0701, DRB1\*0901, DRB4\*0101, DRB5\*0101.

HXB2 Location gp160 (210–218) Author Location Env (186–194 1035) Epitope FEPIPIHYC Subtype B

Immunogen vaccine

Vector/Type: vaccinia prime with gp120 boost Strain: B clade 1035 HIV component: Env Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (Class II I Ab)

Assay type proliferation, T-cell Elispot

Keywords epitope processing, vaccine-induced epitopes,

escape, TCR usage

References Zhan et al. 2003

- A very narrow Th response was stimulated in C57BL/6 mice vaccinated with vaccinia expressed HIV-1 env clone 1035. Five of seven different Th hybridomas isolated from five immunized mice immunized reacted with the peptide PKVSFEPIPIHY-CAP, located in the C2 region of gp120. TCR Vβ usage indicated each of the clones was unique. Splenic populations from other C57BL/6 mice immunized with 1035 env confirmed that the gp120 specific T-helper response was focused on the PKVSFEPIPIHYCAP peptide. The authors suggest the protein structural context may contribute to the immunodominance of this peptide.
- The minimal epitope was mapped for one of the hybridomas, and was FEPIPIHYC.
- The natural variant, fDpipihyc, did not stimulate a response in three of the hybridomas.

HXB2 Location gp160 (210–223)

**Author Location** gp120 (215–228)

Epitope FEPIPIHYCAFPGF

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>bxk</sup>)

References Sastry & Arlinghaus 1991

 Peptides induced T-cell proliferative response to immunizing peptide and to gp160.

**HXB2 Location** gp160 (212–231)

**Author Location** gp120 (221–240 W6.ID)

Epitope PIPIHYCAPAGFAILKCNNK

Immunogen vaccine

Vector/Type: protein Strain: B clade W61D HIV component: gp120 Adjuvant: MPL-SE

adjuvant, QS21

Species (MHC) human

References Jones et al. 1999

- An HIV seronegative volunteer was vaccinated with rgp120 and a QS21/MPL adjuvant and HIV-1 specific T-cell lines were isolated.
- Two T-cell lines react specifically with this peptide.

 $\textbf{HXB2 Location} \hspace{0.1cm} gp160 \hspace{0.1cm} (212\text{--}231)$ 

Author Location gp120 (212–231 IIIB)

Epitope PIPIHYCAPAGFAILKCNNK?

Epitope name D4

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 3/15 responders recognized this peptide, average SI = 4.2.

HXB2 Location gp160 (214–220) Author Location Env (1007) Epitope PIHYCAP

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the C2 region of 1007 (US, clade B) and the Vβ usage of the TCR was not determined.
- The epitope described here is the region of overlap of two 15 mers that were both able to stimulate IL-2 production from the hybridoma (PKVSFEPIPIHYCAP and PIHYCAPAGFAILKC)
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (215–225) **Author Location** Env (1007) Epitope IHYCAPAGFAI

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the C2 region of 1007 (US, clade B) and the  $V\beta$  usage of the TCR was not determined.
- The epitope described here is the region of overlap of two 15 mers that were both able to stimulate IL-2 production from the hybridoma (EPIPIHYCAPAGFAI and IHYCAPAGFAILKCN)
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (216–225)

Author Location Env (UG92005)

Epitope HYCAPAGFAI

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

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**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

 This epitope is located in the C2 region of UG92005 (UG, clade D) and Vβ usage of its TCR was not determined.

- The epitope described here is the region of overlap of two 15 mers that were both able to stimulate IL-2 production from the hybridoma (EPIPIHYCAPAGFAI and HYCAPAG-FAILKCND)
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and  $V\beta$  usage was determined.
- · Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (220-234) Author Location gp120 (225–240 SF2) Epitope PAGFAILKCNNKTFN

Immunogen in vitro stimulation or selection

Species (MHC)

References Manca et al. 1993

- T-cell line derived from unprimed, uninfected individual.
- Responds to APC pulsed with either synthetic peptide or gp120.
- Human MAbs 448-D and 450-D enhance APC gp120 uptake and presentation.

HXB2 Location gp160 (220-234) Author Location gp120 (IIIB) **Epitope** PAGFAILKCNNKTFN

Epitope name pep24 Immunogen vaccine

Vector/Type: Streptococcus gordonii HIV

component: gp120

Species (MHC) human

Keywords immunodominance References Pozzi et al. 1994

• This previously described immunodominant Th cell epitope was fused to the streptococcal surface protein M6 (emm-6.1), for expression on the surface of the bacterium Streptococcus gordonii.

• Recombinant bacteria showed efficient MHC class II mediated presentation of gp120 to T-cells by stimulation of a proliferative response in a human T cell clone specific for pep24.

HXB2 Location gp160 (220-235) Author Location gp120 (IIIB)

Epitope PAGFAILKCNNKTFNY

Immunogen in vitro stimulation or selection

Species (MHC) human (DR2)

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in
- Peptide priming does not always induce T-cells that recognize whole protein.
- gp120 priming induced T-cells that recognize this peptide.

**HXB2 Location** gp160 (220–235) Author Location gp120 (220-235 HXB2) **Epitope** PAGFAILKCNNKTFNY

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DR2)

Keywords escape

References Guzman et al. 1998

• Listeria monocytogenes, an intracellular pathogen which is ingested by macrophages and can escape from the phagosome to replicate in the cytoplasm, was used successfully as carrier to deliver this gp120 epitope to CD4+ T-cells.

HXB2 Location gp160 (220–235) Author Location gp120 (191–205 HXB2)

Epitope PAGFAILKCNNKTFNY

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DR2)

References Fenoglio et al. 1999

- gp120 pep24 epitope exhibited antagonistic activity against proliferation of gp120-specific T-cells when flanked by unrelated amino acid sequence.
- The glutathione S-transferase (GST)-peptide system can be used to display peptides; antigenicity was maintained when this peptide was expressed at the C-term end, but antagonism resulted when this peptide was expressed at the N-term end.

HXB2 Location gp160 (222–241)

Author Location gp120 (222-241 IIIB)

Epitope GFAILKCNNKTFNGTGPCTN?

Epitope name D5

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.

- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 2/15 responders recognized this peptide, average SI = 4.8.

HXB2 Location gp160 (223-231)

Author Location gp120 (194–202 HXB2)

**Epitope** FAILKCNNK

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DR2, DR6) References Manca et al. 1996

- Epitope was the minimal stimulatory sequence defined for two Th lines stimulated in vitro.
- One Th line was stimulated by gp120, one by a Glutathione-Stransferase (GST)-peptide fusion.
- Alanine substitutions at position 914, 196, and 202 abrogated activity for the GST-peptide stimulated line, but not for a gp120 stimulated line.
- · Constructs combining GST and the PAGFAILKCNNKTFNY gp120 peptide at the C-term end of GST stimulated Th cells but not at the N-term end.

HXB2 Location gp160 (223-231)

Author Location gp120 (194–202 HXB2)

**Epitope** FAILKCNNK

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DR2, DR6)

References Manca et al. 1996

- · Epitope was the minimal stimulatory sequence defined for two Th lines stimulated in vitro.
- One Th line was stimulated by p66, one by a Glutathione-Stransferase (GST)-peptide fusion protein.
- Alanine substitutions at position 914, 196, and 202 abrogated activity for the GST-peptide stimulated line, but not for a gp120 stimulated line.
- Constructs linking GST to the PAGFAILKCNNKTFNY gp120 peptide at the C-term end of GST stimulated Th cells, constructs linking at the N-term end did not.
- The C and N termini of GST are not intrinsically permissive or non-permissive, presentation is epitope specific (see SSTVNDIQKLV for contrast)

HXB2 Location gp160 (223-231)

Author Location gp120 (237–245 SF2, HXB2)

Epitope FAILKCNNK

Immunogen

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords subtype comparisons, immunodominance References Fenoglio et al. 2000

- This peptide is an immunodominant Th epitope in BALB/c mice.
- Substitutions in positions 237, 241, 243, 244 with Ala all cause reduced recognition.
- · Most natural analogs they tested did not cross-react, including peptides based on clade A, B, C, D, E and O sequences.
- Position 237 and 244 when substituted with Ala cause an antagonistic response and the natural analogues of this epitope to loose antigenicity.

nistic response.

HXB2 Location gp160 (223-231)

Author Location gp120 (238–246 HXB2)

Epitope FAILKCNNK

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human

Keywords TCR usage

References Li Pira et al. 1998

- · Clonal heterogeneity was broad for a recall response to tetanus toxoid or PPD, but oligoclonal to primary HIV antigens, dominated in this case by TCR  $V\beta$  22 usage.
- Donor of PBMC that recognized this epitope had HLA-DR alleles 2 and 6.
- The only (detected) immunogenic variant of this epitope was derived from strain NOF (YAILKCNNK)

**HXB2 Location** gp160 (230–245)

Author Location gp120 (IIIB)

**Epitope** NKTFNGKGPCTNVSTY

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro.
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (232–251)

Author Location gp120 (232-251 IIIB)

Epitope TFNGTGPCTNVSTVQCTHGI?

Epitope name E1

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- · After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 3/15 responders recognized this peptide, average SI = 3.9.

HXB2 Location gp160 (235–247)

**Author Location** gp120 (240–252)

**Epitope** GTGPCTNVSTVQC

Immunogen vaccine

Vector/Type: peptide

Species (MHC) macaque

References Nehete et al. 1993

Synthetic peptide derived from conserved region of the HIV-1 envelope that stimulates a proliferative response in mice.

munized rhesus monkeys, with a weak transient response in the other two.

HXB2 Location gp160 (238-257) Author Location gp120 (240–249 89.6) Epitope PCTNVSTVQCTHGIRPVVST

Epitope name Peptide 22 Immunogen vaccine

> Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse Donor MHC H-2d

Keywords epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was highly reactive in 6/10 BALB/c mice tested, but not in any (0/10) CBA/J mice.

HXB2 Location gp160 (240-255) Author Location gp120 (IIIB) **Epitope** TNVSTVQCTHGRPIY

**Immunogen** in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

• Peptide stimulation of PBMC from non-infected individuals in vitro.

HXB2 Location gp160 (242-261) Author Location gp120 (242–261 IIIB) Epitope VSTVQCTHGIRPVVSTQLLL Immunogen SHIV infection

Species (MHC) macaque (DRB1\*0406) References Lekutis & Letvin 1997

• A novel C2 region Th epitope was described in SHIV-89.6 infected Macaca mulatta.

HXB2 Location gp160 (242–261) Author Location gp120 (242–261 IIIB) Epitope VSTVQCTHGIRPVVSTQLLL? Epitope name E2

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

• Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.

- Proliferative response to this peptide was observed in 1/3 imspecific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
  - IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
  - 1/15 responders recognized this peptide, SI = 3.4.

HXB2 Location gp160 (244-266)

**Author Location** Env

Epitope TVQCTHGIRPVVSTQLLLNGSLA Epitope name HIV\_env\_DRB0101\_11

Subtype M

Immunogen HIV-1 infection

Species (MHC) human

Country United States. Assay type T-cell Elispot

Keywords computational epitope prediction

References De Groot et al. 2004

- Bioinformatic tools (EpiMatrix and Conservatrix) were employed to predict 9 highly conserved and promiscuous class II restricted T helper epitopes from HIV-1 env sequences for a cross-clade polyepitope vaccine. 10/34 HIV-1 infected patients induced Th1 immune responses to 8/9 study peptides.
- 1/34 subjects tested responded to this peptide.
- Class II peptides containing putative epitopes were selected to favor DRB0101 presentation, as there is a transgenic mouse model for this allele for future vaccine studies. The highest scoring DR B0101 sequence of this peptide was RPVVSTQL.

HXB2 Location gp160 (246–268)

**Author Location** Env (438–460)

Epitope OCTHGIRPVVSTOLLLNGSLAEE

Epitope name HIV\_env\_DRB0101\_02

Subtype M

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assav type T-cell Elispot

**Keywords** computational epitope prediction

References De Groot et al. 2004

- Bioinformatic tools (EpiMatrix and Conservatrix) were employed to predict 9 highly conserved and promiscuous class II restricted T helper epitopes from HIV-1 env sequences for a cross-clade polyepitope vaccine. 10/34 HIV-1 infected patients induced Th1 immune responses to 8/9 study peptides.
- 1/34 subjects tested responded to this peptide.
- Class II peptides containing putative epitopes were selected to favor DRB0101 presentation, as there is a transgenic mouse model for this allele for future vaccine studies. The highest scoring DRB0101 sequence within this peptide was PVVSTQLLL.

HXB2 Location gp160 (250-265)

Author Location gp120 (IIIB)

Epitope GIRPIVSTQLLLNGSC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

Peptide stimulation of PBMC from non-infected individuals in vitro.

• Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (252-271) Author Location gp120 (252–271 IIIB) Epitope RPVVSTQLLLNGSLAEEEVV?

gp160 Helper/CD4+ T-cell epitopes

Epitope name E3 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- · After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 1/15 responders recognized this peptide, average SI = 7.4.

HXB2 Location gp160 (262–281) Author Location gp120 (262–281 IIIB) **Epitope** NGSLAEEEVVIRSVNFTDNA?

Epitope name E4 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 2/15 responders recognized this peptide, average SI = 3.1.

HXB2 Location gp160 (264-287) **Author Location** gp120 (269–292 NL43)

Epitope SLAEEEVVIRSANFTDNAKTIIVQ

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: gp120, gp160

Species (MHC) human

References Sitz et al. 1999

- There was a great breadth of proliferative response to env peptides in 19 HIV-1 infected rgp160 and 17 HIV-1 infected rgp120 vaccine recipients.
- 50% of vaccinees had a stimulation index of greater than 5 to this peptide.

HXB2 Location gp160 (269-283) **Author Location** gp120 (269–283 IIIB, B10) **Epitope** EVVIRSANFTDNAKT

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (269-291)

**Author Location** Env

Epitope EVVIRSENFTNNAKTIIVQLNES

Epitope name HIV\_env\_DRB0101\_7

Subtype M

Immunogen HIV-1 infection

Species (MHC) human

Country United States. Assay type T-cell Elispot

Keywords computational epitope prediction

References De Groot et al. 2004

- Bioinformatic tools (EpiMatrix and Conservatrix) were employed to predict 9 highly conserved and promiscuous class II restricted T helper epitopes from HIV-1 env sequences for a cross-clade polyepitope vaccine. 10/34 HIV-1 infected patients induced Th1 immune responses to 8/9 study peptides.
- 1/34 subjects tested responded to this peptide.
- Class II peptides containing putative epitopes were selected to favor DRB0101 presentation, as there is a transgenic mouse model for this allele for future vaccine studies. The highest scoring DRB0101 sequence of this peptide was NFTNNAKTI.

HXB2 Location gp160 (270–285) Author Location gp120 (IIIB)

Epitope VVIRSDNFTNNAKTIC

**Immunogen** in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (272-291) Author Location gp120 (272–291 IIIB)

Epitope IRSVNFTDNAKTIIVQLNTS?

Epitope name E5 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords immunodominance

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.

- 161), C3 (aa 152-171), C5 (aa 172-191), E5 (aa 272-291) and G4 (aa 380-393). The first three were in or near V2, the other two were proximal to the V3 and V4 loops.
- 4/15 responders recognized this immunodominant peptide, average SI = 5.0.

HXB2 Location gp160 (274–288)

**Author Location** gp120 (274–288 IIIB, B10)

Epitope SANFTDNAKTIIVQL Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (274–296)

Author Location Env

Epitope SENFTNNAKIIIVOLNESVVINV

Epitope name HIV\_env\_DRB0101\_5

Subtype M

Immunogen HIV-1 infection

Species (MHC) human

Country United States. Assav type T-cell Elispot

**Keywords** computational epitope prediction

References De Groot et al. 2004

- · Bioinformatic tools (EpiMatrix and Conservatrix) were employed to select 9 highly conserved and promiscuous class II restricted T helper epitopes from HIV-1 env sequences for a cross-clade vaccine. 10/34 HIV-1 infected patients induced Th1 immune responses to the 9 study peptides.
- 1/26 subjects tested responded to this peptide.
- Class II peptides containing putative epitopes were selected to favor DRB0101 presentation, as there is a transgenic mouse model for this allele for future vaccine studies. The highest scoring DRB0101 sequence of this peptide was AKIIIVQLN.

HXB2 Location gp160 (276-295)

Author Location gp120 (MN)

Epitope NFTDNAKTIIVHLNESVQIN

Epitope name NN20 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type Cytokine production, proliferation, CD4 Tcell Elispot - IFNγ, Intracellular cytokine

staining

Keywords acute/early infection

References Malhotra et al. 2003

• 92 acute- or early-HIV infected subjects were tested for Envand Gag-specific Th responses. There was an overall low probability of detecting HIV-1-specific Th responses, Env responses were rare and (5%) and only detected very early in infection, probably reflecting a low frequency of response and impaired lymophoproliferative capacity, not viral escape. Gagspecific Th responses were observed more frequently (16%), were found throughout the acute and early infection phases, and predominated after ARV therapy.

• Five peptides were recognized most frequently: C2 (aa 142- • This epitope is one of six Env-specific Th responses detected in one patient 11 days post-infection that were studied in detail. These Th reponses were not detected at subsequent time points, despite the Env specific Th cells being maintained as detected by TCR clonal transcripts, and the epitope sequence being maintained in Env.

HXB2 Location gp160 (280–296)

Author Location gp120 (IIIB)

Epitope NAKTIIVQLNESVAIC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (288–307)

**Author Location** gp120 (290–309 89.6)

Epitope LNESVVINCTRPNNNTRRRL

Epitope name Peptide 27 Immunogen vaccine

> Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse

Donor MHC H-2k, H2-d

**Keywords** epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was reactive in only 1/10 BALB/c mice tested, but reacted in 8/10 CBA/J mice.

HXB2 Location gp160 (289–297)

Author Location gp120 (292–300 SF2)

Epitope NESVAINCT Immunogen vaccine

Vector/Type: protein Strain: B clade SF2

HIV component: gp120

Species (MHC) human

References Botarelli et al. 1991

• A non-glycosylated form of SF2 gp120, env 2-3, was used as an immunogen - 20% of T-cell clones do not recognize the glycosylated form.

**HXB2 Location** gp160 (290–306)

Author Location gp120 (296–312 LAI)

**Epitope** SVVEINCTRPNNNTRKS

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

HXB2 Location gp160 (290-314)

Author Location Env

Epitope ESVVINCTRPNNNTRRSIHIGPG

Epitope name HIV\_env\_DRB0101\_14

Subtype M

Immunogen HIV-1 infection

Species (MHC) human

Country United States. Assay type T-cell Elispot

Keywords computational epitope prediction

References De Groot et al. 2004

- Bioinformatic tools (EpiMatrix and Conservatrix) were employed to predict 9 highly conserved and promiscuous class II restricted T helper epitopes from HIV-1 env sequences for a cross-clade polyepitope vaccine. 10/34 HIV-1 infected patients induced Th1 immune responses to 8/9 study peptides.
- 1/34 subjects tested responded to this peptide.
- Class II peptides containing putative epitopes were selected to favor DRB0101 presentation, as there is a transgenic mouse model for this allele for future vaccine studies. The highest scoring DR B0101 sequence of this peptide was TRPNNNTRR.

**HXB2 Location** gp160 (292–310)

Author Location gp120 (292–310 IIIB)

Epitope VEINCTRPNNNTRKRIRIQ?

Epitope name F1

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- Only 1/15 responders recognized this peptide, but it had the highest SI in the study of 9.9.

HXB2 Location gp160 (296–307)

Author Location gp120 (301–324 RF)

**Epitope** CTRPNNNTRKSI **Immunogen** HIV-1 infection

Species (MHC)

Keywords epitope processing

References de Lorimier et al. 1994

Proton NMR spectroscopy was employed to analyze the solution conformation of a hybrid peptide, T1-SP10RF, in order to better understand the immunogenicity of its' T helper (KQI-INMWQEVGKAMYA, CTRPNNNTRKSI), CTL (SITKGP-GRVIYATG) and B-cell epitopes (RKSITKGPGRVIYATG).

 This epitope embedded in the T1-SP10RF peptide does not form a helical amphipathic conformation. It lacks random-coil conformations, and this may make a peptide less susceptible to complete proteolytic degredation and be favored within epitopes.

**HXB2 Location** gp160 (296–314)

Author Location gp120 (303–321 IIIB)

Epitope CTRPNNNTRKSIRIQRGPG(Y)

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

Species (MHC) goat

References Palker et al. 1989

• Goats were immunized with peptides containing V3 typespecific neutralizing determinants coupled to T1.

HXB2 Location gp160 (297–321)

Author Location gp120 (302-324 MN)

Epitope TRPNYNKRKRIHIGPGRAFYTTK

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN

HIV component: V3

**Species (MHC)** mouse (H-2<sup>d</sup>)

**References** Oscherwitz *et al.* 1999b

- Epitope presented as a tandem repeat (eight copies) elicits stronger B-cell and T-cell responses than the epitope presented as a single copy.
- This study indicates that the increased response was not due to neodeterminants created at the junction of the peptides, but rather due to an epitope density effect, increased immunogenicity through a high ratio of epitope to protein.

HXB2 Location gp160 (297-330)

**Author Location** Env (303–335 BX08)

Epitope TRPNNNTRKSIHIGPGRAFYATGEIIGDIRQAH

Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I trial.
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees – 6/10 reacted to this peptide.
- 9/12 tested mounted a CTL responses to at least one of the six peptides, each of the six peptides elicited a CTL response in at least one individual – this peptide was particularly immunogenic, eliciting a CTL response in five vaccinees.
- None of the 12 tested had an IgG response to gp120 or gp160 and vaccinees could be differentiated from HIV-1 seropositive individuals with a commercial HIV detection kit – no neutralizing antibodies were observed.

**HXB2 Location** gp160 (298–307)

**Author Location** Env (UG92005)

Epitope RPYNNTRKGI

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the V3 region of UG92005 (UG, clade D) and was recognized by a hybridoma with Vβ usage not determined.
- The epitope described here is the region of overlap of two 15 mers that were both able to stimulate IL-2 production from the hybridoma (TINCTRPYNNTRKGI and RPYNNTRKGI-HIGPG)
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (298–319) **Author Location** gp120 (300–319 89.6)

Epitope RPNNNTRRRLSIGPGRAFYA

**Epitope name** Peptide 28 **Immunogen** vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse

**Donor MHC** H-2k, H2-d **Keywords** epitope processing, immunodo

**Keywords** epitope processing, immunodominance **References** Dai *et al.* 2001

 Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.

• This peptide was reactive in 7/10 BALB/c mice tested, and in 5/10 CBA/J mice.

 $\textbf{HXB2 Location} \hspace{0.1cm} \texttt{gp160} \hspace{0.1cm} (301\text{--}325)$ 

Author Location gp120 (IIIB)

Epitope NNTRKSIRIQRGPGRAFVTIGKIGN

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB HIV component: Env, Rev Adjuvant: QS21

Species (MHC) mouse Keywords Th1

References Sasaki et al. 1998

- The env response is what is being sought, but co-expression of rev is required.
- Intramuscular versus nasal vaccination with DNA vaccine with a QS-21 adjuvant was studied.
- QS-21 enhanced the IgG2a response mediated via Th1 cytokines IFN $\gamma$  and IL-2 and delayed type hypersensitivity (DTH) in response to the V3 peptide was measured by a foot pad swelling test Sasaki *et al.* [1998]

**HXB2 Location** gp160 (302–315)

Author Location gp120 (307–322 IIIB)

Epitope NTRKSIRIQRGPGR

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

HIV component: V3

Species (MHC) mouse

References Goodman-Snitkoff et al. 1990

 Identification of putative Th epitopes that can stimulate an antibody response in peptide-immunized mice.

**HXB2 Location** gp160 (302–321)

Author Location gp120 (302-321 IIIB)

Epitope NTRKRIRIQRGPGRAFVTIG?

**Epitope name** F2

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 1/15 responders recognized this peptide, SI = 5.6.

HXB2 Location gp160 (302–327)

**Author Location** gp120 (307–332 MN)

 ${\bf Epitope} \ \ {\tt NKRKRIHIGPGRAFYTTKNIIGTIR}$ 

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN HIV component: V3 Adjuvant: Montanide (ISA 51)

Species (MHC) mouse

References Anderson et al. 2001

Hypervariable epitope constructs (HECs) are degenerative peptide cocktails that are made in a single peptide synthesis reaction. Vaccination with a V3 degenerative peptide cocktail containing 64 distinct peptides, NTRK-[SR]-I-[HR]-IGPG-[RQ]-AFY-[AT]-TG-[DE]-IG-[DN]-IRQ, elicited broader and more durable Th responses than the MN V3 peptide alone in BALB/c mice immunized and boosted with V3 peptides, although the MN peptide elicited a transient MN-specific V3 response.

**HXB2 Location** gp160 (305–321) **Author Location** gp120 (312–329)

Epitope (CG)KSIRIQRGPGRAFVTIG

Immunogen HIV-1 infection

Species (MHC) human

References Adams et al. 1997

• Used as positive control in study examining T-cell response to four p24 Gag peptides.

HXB2 Location gp160 (308–319)
Author Location gp120 (subtype C)
Epitope (CKR)KIHIGPGQAFYT

Subtype C

Immunogen HIV-1 infection

**Species (MHC)** mouse (H-2<sup>b</sup>, H-2<sup>d</sup>, H-2<sup>k</sup>, H-2<sup>s</sup>)

Keywords Th1

References Ahluwalia et al. 1997

 A V3 loop peptide modified to resemble an Indian form (GPGQ) was incorporated into ISCOMS (immune stimulating complexes) or liposomes, and used to immunize mice – the IgG2a/IgG2b Ab response was enhanced by the presentation in the ISCOM suggestive of a Th1 response.

HXB2 Location gp160 (308–321) Author Location gp120 (MN) Epitope RIHIGPGRAFYTTK

Epitope name SP10 Immunogen vaccine

Vector/Type: peptide Strain: B clade MN HIV component: V3

**Species (MHC)** mouse (H-2<sup>d</sup>) **References** Klinman *et al.* 1995

Hybrid T1-V3 peptide activates IL-4 and IL-6 in a dose dependent manner.

• 10-mer from V3 contributes to this response.

HXB2 Location gp160 (308–322) Author Location gp120 (315–329 IIIB) Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen HIV-1 infection
Species (MHC) human (DR)
References Baier et al. 1995

 Linked HIV-1 T1 and P18 peptides to anti-HLA-DR and IgD Fab fragments to enhance uptake by antigen presenting cells thus increase immunogenicity.

**HXB2 Location** gp160 (308–322) **Author Location** gp120 (315–329 IIIB)

**Epitope** RIQRGPGRAFVTIGK **Epitope name** P18

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species (MHC)** mouse (H-2 A<sup>d</sup>) **References** Takahashi *et al.* 1990

 Induces both class II restricted CD4+ Th cells, and class I restricted CD8+ CTL.

HXB2 Location gp160 (308–322)

Author Location gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen peptide-HLA interaction

**Species (MHC)** mouse (H-2 I-A<sup>d</sup>) **References** Takeshita *et al.* 1995

 Binds Class II H-2 I-A<sup>d</sup> requiring riqrgPgRaFvti, and Class I H-2 D<sup>d</sup>, requiring iGPgRaFvtI.

HXB2 Location gp160 (308–322) Author Location Env (IIIB)

Epitope RIQRGPRAFVTIGK

Epitope name P18 Immunogen vaccine

*Vector/Type:* DNA with CMV promotor *Strain:* B clade IIIB *HIV component:* gp160, Rev *Adjuvant:* MIP-1 $\alpha$ 

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1

References Lu et al. 1999

- MIP-1a expression plasmid co-inoculated with a DNA vaccine consisting of HIV-1 pCMV160IIIB and pcREV enhanced the HIV-specific T-cell immune response as measured by a CTL test against using V3 peptide pulsed targets, and a DTH test to V3 peptide.
- The IgG1/IgG2a response was lowered with co-inoculation of MIP-1 alpha, suggesting it preferentially elicits a Th1 response.

HXB2 Location gp160 (308–322) Author Location gp120 (308–322 IIIB) Epitope RIHIGPGRAFYTTKN

Immunogen

Species (MHC) human

References Furci et al. 1997

- 9/11 exposed-uninfected individuals in this study had a proliferative response to a C5 peptide, but only 1/11 exposeduninfected individuals recognized this peptide.
- 1/18 unexposed-uninfected controls could recognize this peptide.
- Erroneously documented as IIIB sequence most likely MN peptide.

HXB2 Location gp160 (308–322)

**Author Location** gp120 (315–329 IIIB) **Epitope** RIQRGPGRAFVTIGK

Epitope name P18

Immunogen vaccine

Vector/Type: peptide

Species (MHC) macaque

References Nehete et al. 1993

- Synthetic peptide derived from conserved region of the HIV-1 envelope that stimulates a proliferative response in mice.
- Despite the proliferative response to this peptide in mice and humans, no response was observed in 3 rhesus monkeys.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen HIV-1 infection

Species (MHC) human

Keywords responses in children, Th1, Th2

References Wasik et al. 1997

- The breadth and intensity of the CTL response and the type of Th response was studied in seven rapidly progressing HIV-1 + infants.
- IL-2 and  $\gamma$  IFN production from Th1 cells correlated with the CTLp frequency against HIV-1 Gag, Env, Nef and Pol.
- IL-4 production from Th2 cells was inversely correlated with the CTLp frequency.
- The HIV-1 + children with strong CTL responses had levels of anti-CD3 MAb induction of Th1 cells comparable to uninfected children.
- The children that did not mount a good CTL response had dramatically decreased numbers of Th1 relative to Th2 cells.

HXB2 Location gp160 (308-322)

Author Location gp120 (315-329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** responses in children, kinetics, Th1

References Wasik et al. 2000

- Th responses measured by IL-2 responses to P18 and T1 in HIV-1 infected infants were undetectable at less than 1 month of age, and remained low in children with AIDS symptoms, but increased with age in children with slowly progressive disease.
- The kinetics and intensity of the CTL activity during the first year of life was related to the child's ability to make Th1 responses.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (315-329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen

Species (MHC) human

References Pinto et al. 1995

 CTL activity analyzed in parallel with Th reactivity in exposed but uninfected health care workers.

HXB2 Location gp160 (308–322)

Author Location gp120 (315–329 MN)

Epitope RIHIGPGRAFYTTKN

Epitope name P18

Immunogen

Species (MHC) human

References Pinto et al. 1995

 CTL activity analyzed in parallel with Th reactivity in exposed but uninfected health care workers.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (315-329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1989

• IL-2 production detection of Th lymphocytes from asymptomatic HIV-positive individuals.

**HXB2 Location** gp160 (308–322)

**Author Location** gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1991a

 Peptides stimulate Th cell function and CTL activity in similar patient populations.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp160

Species (MHC) human

References Clerici et al. 1991b

• Immunizing uninfected individuals with rgp160 results in stronger Th response than does natural infection.

HXB2 Location gp160 (308-322)

Author Location gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen

Species (MHC) human

**References** Clerici *et al.* 1992

• Cell-mediated immune response to HIV-1 peptides in HIV-1 exposed seronegative men.

**HXB2 Location** gp160 (308–322)

Author Location gp120 (315-329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1997

used in a study of the influence of pentoxifylline on HIV specific T-cells.

**HXB2 Location** gp160 (308–322) **Author Location** gp120 (MN)

Epitope RIHIGPGRAFYTTKN

Immunogen

Species (MHC) human

References Clerici et al. 1992

• Epitope P18 MN: Cell-mediated immune response to HIV-1 peptides in HIV-1 exposed seronegative men. • Measurable HIV specific T help responses elicited in the immunologically immature newborn, possibly in response to *in* 

**HXB2 Location** gp160 (308–322)

Author Location gp160 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human

**Keywords** immunodominance **References** Wasik *et al.* 1999

- IL-2 responses associated with beta-chemokine expression were detectable at birth in the majority of uninfected infants born to HIV+ mothers, declining by age 6 months.
- In both uninfected and infected infants of HIV-positive mothers, responses to the T1 peptide (KQIINMWQEVGKAMYA) were more frequent than responses to P18.
- T1 is a highly conserved epitope, whereas P18 has a higher mutation rate due to its location in the immunodominant V3 loop region.

**HXB2 Location** gp160 (308–322)

**Author Location** gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen HIV-1 infection

Species (MHC) human

References Kaul et al. 1999

- Kenyan sex workers that remained seronegative were found to frequently have HIV-env peptide specific Th responses detected by an IL-2 assay (11/20 cases) and mucosal genital tract anti-HIV IgA (16/21 cases)
- Helper epitopes used in this study were noted to be previously described Clerici et al. [1989], and were not explicitly described in Kaul et al. [1999]

 $\textbf{HXB2 Location} \ \ gp160 \ (308-322)$ 

Author Location gp120 (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human

Keywords subtype comparisons, responses in children,

mother-to-infant transmission

References Kuhn et al. 2001a

- In a S. African perinatal transmission study, 33% (33/86) of cord blood samples from infants with seropositive mothers produced T-helper responses (measured by a bioassay measuring IL2 production in a murine cell line and confirmed with a proliferation assay) against a peptide cocktail containing The epitopes P18 MN, P18 IIIB, T1, T2, and TH4.
- The mothers were predominantly infected with subtype C, but the T help response was detectable in a number of cord blood samples despite using peptides based on B subtype reagents.

- 3/33 infants with cord blood T help responses to Env were infected *in utero*, 2/33 were lost to follow up, and 28/33 were not infected. 6/53 of the infants with cord blood that was unresponsive to Env peptide stimulation were infected before delivery, and 8/47 contracted HIV intrapartum or via breast-feeding.
- Measurable HIV specific T help responses elicited in the immunologically immature newborn, possibly in response to in utero exposure, are associated with a protective natural immunity that helps block mother-infant transmission of HIV-1.

HXB2 Location gp160 (308–322)

Author Location gp120 (315-329 MN)

Epitope RIHIGPGRAFYTTKN

Epitope name P18

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human

Keywords subtype comparisons, responses in children,

mother-to-infant transmission

References Kuhn et al. 2001a

- In a S. African perinatal transmission study, 33% (33/86) of cord blood samples from infants with seropositive mothers produced T-helper responses (measured by a bioassay measuring IL2 production in a murine cell line and confirmed with a proliferation assay) against a peptide cocktail containing Th epitopes P18 MN, P18 IIIB, T1, T2, and TH4.
- The mothers were predominantly infected with subtype C, but the T help response was detectable in a number of cord blood samples despite using peptides based on B subtype reagents.
- 3/33 infants with cord blood T help responses to Env were infected *in utero*, 2/33 were lost to follow up, and 28/33 were not infected. 6/53 of the infants with cord blood that was unresponsive to Env peptide stimulation were infected before delivery, and 8/47 contracted HIV intrapartum or via breast-feeding.
- Measurable HIV specific T help responses elicited in the immunologically immature newborn, possibly in response to in utero exposure, are associated with a protective natural immunity that helps block mother-infant transmission of HIV-1.

**HXB2 Location** gp160 (308–322)

Author Location Env (315–329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18IIIB

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC)

Assay type Cytokine production

Keywords mother-to-infant transmission

References Clerici et al. 1993a

- Cord blood samples in 8/23 infants with HIV+ mothers showed IL-2 production in response to peptides from HIV-1 gp1260, demonstrating that Th cell priming to HIV env determinants can occur in utero. HIV-specific Th immunity in utero may be protective, as none of the 8 with HIV-1 specific Th activity became infected, while 3/15 infants with no detectable Th activited were infected.
- PBL from 10/21 of the mother showed HIV-1 specific Th activity through IL-2 production in response to Env peptides.

HXB2 Location gp160 (308–322)

**Author Location** Env (MN)

Epitope RIHIGPGRAFYTTKN

Epitope name P18MN

Immunogen HIV-1 infection, HIV-1 exposed seronegative Species (MHC)

Assay type Cytokine production

Keywords mother-to-infant transmission

References Clerici et al. 1993a

- Cord blood samples in 8/23 infants with HIV+ mothers showed IL-2 production in response to peptides from HIV-1 gp1260, demonstrating that Th cell priming to HIV env determinants can occur in utero. HIV-specific Th immunity in utero may be protective, as none of the 8 with HIV-1 specific Th activity became infected, while 3/15 infants with no detectable Th activited were infected.
- PBL from 10/21 of the mother showed HIV-1 specific Th activity through IL-2 production in response to Env peptides.

HXB2 Location gp160 (308-322)

**Author Location** Env (IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18IIIB

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC)

Assay type Cytokine production

References Clerici et al. 1994a

- Six of eight HIV-exposed health care workers had transient HIV-specific T-helper responses after percutaneous exposure to HIV, responding to two or more Env peptides among the set T1, T2, Th4.1, P18IIIB and P18MN. Responses were lost after 12-56 weeks. The HIV-specific Th responses occurred without seroconversion or PCR evidence for infection.
- Six of the eight HIV-exposed individuals responded to two or more peptides, whereas, only one individual exposed to HIV-negative blood responded to one peptide.

HXB2 Location gp160 (308-322)

Author Location Env (MN)

Epitope RIHIGPGRAFYTTKN

Epitope name P18MN

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC)

Assay type Cytokine production

References Clerici et al. 1994a

- Six of eight HIV-exposed health care workers had transient HIV-specific T-helper responses after percutaneous exposure to HIV, responding to two or more Env peptides among the set T1, T2, Th4.1, P18IIIB and P18MN. Responses were lost after 12-56 weeks. The HIV-specific Th responses occurred without seroconversion or PCR evidence for infection.
- Six of the eight HIV-exposed individuals responded to two or more peptides, whereas, only one individual exposed to HIV-negative blood responded to one peptide.

HXB2 Location gp160 (308-322)

**Author Location** HIV-1 (IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18IIIB

Subtype B

Immunogen HIV-1 infection

Species (MHC)

Assay type Cytokine production References Clerici et al. 1994b

• IL-10-specific mRNA was upregulated in PBMC from asymptomatic, HIV-infected (HIV+) patients, and was particularly high in those with severely compromised Th cells function. Th response to HIV peptides in vitro could be restored by IL-10

HXB2 Location gp160 (308-322)

**Author Location** HIV-1 (MN)

Epitope RIHIGPGRAFYTTKN

Epitope name P18MN

Subtype B

Immunogen HIV-1 infection

Species (MHC)

Assay type Cytokine production

References Clerici et al. 1994b

• IL-10-specific mRNA was upregulated in PBMC from asymptomatic, HIV-infected (HIV+) patients, and was particularly high in those with severely compromised Th cells function. Th response to HIV peptides in vitro could be restored by IL-10

HXB2 Location gp160 (308–322)

**Author Location** Env (315–329)

**Epitope** RIHIGPGRAFYTTKN

Epitope name P18 MN

Immunogen HIV-1 infection

Species (MHC) human

Assay type Cytokine production

**Keywords** mother-to-infant transmission

References Kuhn et al. 2001b

- The proliferative responses in cord blood at delivery to a cocktail of HIV Envelope peptides were infrequent (1/41) among infants whose HIV-positive mothers received antiretroviral prophylaxis at delivery to prevent transmission. Responses were 10 times more frequent among infants of HIV-seropositive women who had no antiretroviral treatment (7/29). Reductions of HIV-specific responses occurred despite persistance of detectable HIV RNA in the mothers at delivery.
- The reduction of Th responses in newborns raises the possibility that anti-retrovial exposure during pregnancy may block subsequent immune protection. The authors point two relevant citations that showed ARV exposed infants had a more rapid course of progression (Kuhn et al., JID 182:104 (2000)), but were at no greater risk of infection due to subsequent breast feeding (Ditrame et al., Lancet 354:2050 (1999)).

HXB2 Location gp160 (308–322)

Author Location Env (315-329 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18 IIB

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type proliferation

**Keywords** responses in children, mother-to-infant transmission

References Kuhn et al. 2001b

- T helper proliferative responses in cord blood at delivery to a cocktail of HIV envelope peptides were infrequent (1/41) among infants whose HIV-positive mothers received antiretroviral prophylaxis at delivery to prevent transmission. Responses were 10 times more frequent among infants of HIV-seropositive women who had no antiretroviral treatment (7/29). Reductions of HIV-specific responses occurred despite persistance of detectable HIV RNA in the mothers at delivery.
- The reduction of Th responses in newborns raises the possibility that anti-retrovial exposure during pregnancy may block subsequent immune protection. The authors point two relevant citations that showed ARV exposed infants had a more rapid course of progression (Kuhn *et al.*, JID 182:104 (2000)), but were at no greater risk of infection due to subsequent breast feeding (Ditrame *et al.*, Lancet 354:2050 (1999)).

HXB2 Location gp160 (308–322)

Author Location Env (gp160) (317–331 MN)

Epitope RIHIGPGRAFYTTKN

Epitope name P18 Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Country** South Africa. **Assay type** proliferation

**Keywords** responses in children, variant crossrecognition or cross-neutralization

References Meddows-Taylor et al. 2004

- Viral isolates (gp160) from 16 vertically HIV-1 infected children (8 T-helper cell Env responders and 8 non-responders) were analyzed for variation after comparison with Env peptide sequences (conserved regions T1, T2 and TH4.1, hypervariable regions P18 MN and P181 IIIB) used for in vitro stimulation.
- No correlation between the age, clinical category (mild or severe), HIV-1 viral load and the degree of epitope variation was established.

HXB2 Location gp160 (308-322)

Author Location Env (gp160) (317–331 IIIB)

Epitope RIQRGPGRAFVTIGK

Epitope name P18 Subtype C

Immunogen HIV-1 infection

Species (MHC) human

**Country** South Africa. **Assay type** proliferation

**Keywords** responses in children

References Meddows-Taylor et al. 2004

- Viral isolates (gp160) from 16 vertically HIV-1 infected children (8 T-helper cell Env responders and 8 non-responders) were analyzed for variation after comparison with Env peptide sequences (conserved regions T1, T2 and TH4.1, hypervariable regions P18 MN and P181 IIIB) used for in vitro stimulation.
- No correlation between the age, clinical category (mild or severe), HIV-1 viral load and the degree of epitope variation was established.

**HXB2 Location** gp160 (308–327)

Author Location gp120 (306–325 MN)

Epitope RIHIGPGRAFYTTKNIIGIT

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101)

References Hayball et al. 1997

- Tandem repeated presentation of epitope enhances binding to class II molecule and therefore induction of T-cell proliferation.
- Tandem peptides are thought to enhance proliferation through improved recruiting of CD4 to the activation complex, which can counter-balance gp120's sequestering of CD4 and consequential inhibition of a proliferative response.

HXB2 Location gp160 (309-323)

**Author Location** gp120 (309–323 IIIB, B10)

Epitope EQRGPGRAFVTIGKI

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

**HXB2 Location** gp160 (309–325)

Author Location gp120 (314–330)

Epitope IQRGPGRAFVTIGKIGN

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Caruso *et al.* 1997

- As HIV-1-infected individuals progress to disease, T-cells show reduced ability to proliferate in response to HIV antigen, but retain the ability to express the activation antigens CD25 and CD71.
- The ability to express activation markers in response to HIV is retained, but the response to tetanus toxoid recall antigen is lost.
- This study investigated CD25 and CD71 expression in PBMC from patients at various stages of progression, measuring the response to *in vitro* stimulation by peptide cocktail containing four antigenic Env peptides, or p17 and p24.

**HXB2 Location** gp160 (310–328)

**Author Location** gp120 (310–329 89.6)

Epitope SIGPGRAFYARRNIIGDIRQ

**Epitope name** Peptide 29

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse

Donor MHC H-2k, H2-d

Keywords epitope processing, immunodominance

References Dai et al. 2001

 Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal gen processing and the frequency of immunogenic sequences.

• This peptide was reactive in 2/10 BALB/c mice tested, and in 8/10 CBA/J mice.

HXB2 Location gp160 (311-319)

**Author Location** 

Epitope RGPGRAFVT

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade BH10 HIV component: gp120 Adjuvant: GM-CSF

Species (MHC) mouse

References Barouch et al. 2002

- gp120 encoding DNA co-injected with a plasmid carrying GM-CSF gave meager CD4+ T-cell responses in BALB/c mice relative to bicistronic gp120 and GMCSF cloned into the same vector and expressed from the same promoter.
- The bicistronic gp120/GM-CSF vaccine induced an approximately 10-fold increase of CD4+ T cell proliferative responses to gp120, as well as a significant increase in IL-2, IL-4, IL-10, IFNγ and GM-CSF production, compared to immunization with the monocistronic pVIJ-gp120 with GMCSF. The enhanced proliferative responses were substantiated by CD4+ T-cell Elispot.
- · Both mono and bicistronic DNA vaccines induced similar CTL responses directed against the H-2Dd restricted P18 peptide RGPRAFTVTI in murine splenocytes despite the enhanced proliferative responses.

HXB2 Location gp160 (311-320)

Author Location gp120 (IIIB)

Epitope RGPGPAFVTI

Immunogen vaccine

Vector/Type: DNA with CMV promotor B clade IIIB HIV component: Strain: gp160, Rev Adjuvant: IL-2

Species (MHC) mouse (H-2<sup>d</sup>)

Keywords Th1

References Xin et al. 1998

• Intranasal immunization with IL-2 expression plasmid in addition to DNA vaccine amplifies cellular response to antigen, probably via activation of Th type 1 (Th1) cells.

HXB2 Location gp160 (311-320)

Author Location gp120 (IIIB)

Epitope RGPGPAFVTI

Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade IIIB HIV component: gp160, Rev Adjuvant: IL-15

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1

References Xin et al. 1999

· Intranasal immunization with IL-15 expression plasmid in addition to DNA vaccine increases DTH response and CTL activity to the antigen, and decreases the serum IgG1 to IgG2a ratio, enhancing Th type 1 (Th1) cell-mediated immunity.

segments, suggesting 3-D protein structure influences Th anti- • Expression of IL-2 or IL-15 can enhance Th1 response to the vaccine, but they to do not appear to elicit a synergistic response.

HXB2 Location gp160 (311–320)

Author Location gp120 (IIIB)

Epitope RGPGPAFVTI

Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade IIIB HIV component: gp160, Rev Adjuvant: CD40

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1, Th2

References Ihata et al. 1999

- CD40L expression increases DTH, and Th1-dependent responses based on enhanced IgG2a titers, with no lowering of IgG1 titers.
- Elispot assay indicated co-injection with hCD40L resulted in greater numbers of IFN \( \gamma \) producing Th1cells, as well as increased IL-4 producing Th2 cells.
- Results suggest hCD40L enhance both Th1 and Th2 cells, and such a pattern of induction is unique among adjuvants, as most adjuvants increase either Th1 or Th2.

HXB2 Location gp160 (311–322)

Author Location Env (IIIB)

Epitope RGPGRAFVTIGK

Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade IIIB HIV component: gp160, Rev Adjuvant: GM-CSF

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1, Th2

References Kusakabe et al. 2000

The timing of delivery of the pGM-CSF expression plasmid for intramuscular DNA pCMV160IIIB/REV vaccination impacts the Th response, maximizing Th2 responses when administered 3 days prior to the DNA vaccine, and Th1 responses when administered 3 days after the DNA vaccine.

HXB2 Location gp160 (314–328)

**Author Location** gp120 (314–328 IIIB, B10)

Epitope GRAFVTIGKIGNMRQ

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

**HXB2 Location** gp160 (314–341)

**Author Location** gp120 (319–346 NL43)

Epitope GRAFVTIGKIGNMRQAHCNISRAKWNAT

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: gp120, gp160

Species (MHC) human

References Sitz et al. 1999

There was a great breadth of proliferative response to env peptides in 19 HIV-1 infected rgp160 and 17 HIV-1 infected rgp120 vaccine recipients.

• More than 25% of vaccinees had a stimulation index of greater than 5 to this peptide.

HXB2 Location gp160 (315–328) Author Location Env (UG92005) Epitope RAYYTTNIVGNIRQ

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the V3 region of UG92005 (UG, clade D) and was recognized by two hybridomas with  $V\beta$  usage not determined, but one used  $V\alpha$  8.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (317–331)

**Author Location** gp120 (324–338 IIIB)

Epitope FVTIGKIGNMRQAHC

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup>) **References** Hale *et al.* 1989

 Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

HXB2 Location gp160 (317–331) Author Location gp160 (324–338 IIIB) Epitope FVTIGKIGNMRQAHC Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup>)

References Berzofsky *et al.* 1991b; Berzofsky *et al.* 1991a

• B10.BR (H-2A<sup>k</sup>, E<sup>k</sup>) and B10.D2 (H-2A<sup>d</sup>, E<sup>d</sup>) mice immunized with rec gp160 showed a proliferative response to this peptide.

FVTIGKIGNMRQAHCNISRAKWNNTLKQIDSKL encompasses several murine Th epitopes including FVTIGKIGNM-RQAHC and is referred to as a "multideterminant region" or cluster peptide.

HXB2 Location gp160 (317–336)

Author Location gp120 (321–340 MN)

Epitope YTTKNIIGTIRQAHCNSRA

Epitope name 1987 Subtype B Immunogen vaccine

Vector/Type: DNA, protein Strain: B clade MN HIV component: gp120 Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) guinea pig

**Keywords** vaccine-specific epitope characteristics, Th1 **References** Chattergoon *et al.* 2002

- Hartley guinea pigs were intradermally injected with either recombinant protein or plasmid expressed gp120 and monitored for delayed type hypersensivity (DTH) responses after vaccination, which are related to Th1 T-cell responses. CFA did not augment responses in animals vaccinated with plasmid.
- A total of 7 gp120 peptides elicited a delayed type hypersensivity (DTH) response after vaccination, out of a set of 60 overlapping peptides that spanned gp120. The vaccine delivery system, DNA versus rec protein, resulted in the recognition of distinct peptides.
- 4/5 animals vaccinated with rec gp120 responded by DTH to this peptide, as did 4/6 vaccinated with plasmid gp120 DNA.

**HXB2 Location** gp160 (317–349) **Author Location** gp160 (324–356 IIIB)

Epitope FVTIGKIGNMRQAHCNISRAKWNNTLKQIDSKL

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** human, mouse (H-2<sup>k</sup>, H-2<sup>d</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- FVTIGKIGNMRQAHCNISRAKWNNTLKQIDSKL encompasses several murine Th epitopes and is referred to as a "multideterminant region" or cluster peptide.
- Six multideterminant region cluster peptides were evaluated Th responses in different MHC/HLA backgrounds after vaccination of mice with gp160, or in infected people.
- This cluster peptide elicited proliferative responses in cells from B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>) and B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), but shorter peptides from within this region stimulated H-2<sup>k</sup>, H-2<sup>d</sup>, H-2<sup>b</sup> and H-2<sup>s</sup> responses.
- IL-2 production in response to this peptide was observed in 58%~(21/36) of asymptomatic HIV-infected individuals.

HXB2 Location gp160 (319–338)

Author Location gp120 (320–339 89.6)

Epitope RRNIIGDIROAHCNISRAKW

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse (H-2<sup>k</sup>, H-2<sup>d</sup>) Keywords immunodominance References Dai *et al.* 2001

- Promiscuous immunodominant epitopes in gp120 were mapped by overlapping peptides in CBA/J H-2<sup>k</sup> and BALB/c H-2<sup>d</sup> mice, and all were found to be in the outer domain, proximal to regions of structural disorder indicated by the crystal structure or by sequence divergence.
- This peptide was recognized by 7/10 CBA/J and 7/10 BALB/c mice with SI > 4, averaging 6.3 and 4.8, and is considered to be promiscuously immunodominant.
- Uniquely immunodominant sequences tended to be in the inner domain of the protein.

HXB2 Location gp160 (319–338)

Author Location gp120 (320–339 89.6)

Epitope RRNIIGDIRQAHCNISRAKW

**Epitope name** Peptide 30 **Immunogen** vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse Donor MHC H-2k, H2-d

**Keywords** epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was highly reactive in 7/10 BALB/c mice tested, and in 7/10 CBA/J mice and was considered one of the 3 immunodominant peptides identified that were shared in both mouse strains: RRNIIGDIRQAHCNISRAKW, NNTLQQIVIKLREK-FRNKTI, GTNGTEGNDIITLQCRIKQI.

HXB2 Location gp160 (321–336) Author Location gp120 (IIIB)

Epitope RIIGDIRKAHCNISRY

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro
- Peptide priming does not always induce T-cells that recognize whole protein.

**HXB2 Location** gp160 (322–336) **Author Location** Env (1007) Epitope IIGDIRQAHCNISRE

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the V3 region of 1007 (US, clade B) and was recognized by three hybridomas with Vβ usage Vβ 6 and not determined.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (322–336) Author Location Env (UG92005) Epitope IVGNIRQAHCNVSKA

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

• This epitope is located in the V3 region of UG92005 (UG, clade D) and was recognized by three hybridomas with  $V\beta$  usage  $V\beta$  6, 8.1, and not determined.

- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (322–336) Author Location Env (UG92005) Epitope IVGNIRQAHCNVSKA

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the V3 region of UG92005 (UG, clade D) and was recognized by three hybridomas with  $V\beta$  usage  $V\beta$  6, 8.1, and not determined.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA transfected L cells as targets and  $V\beta$  usage was determined.

- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (322–341) **Author Location** gp120 (322–341 IIIB)

Epitope KIGNMRQAHCNISRAKWNNT?

Epitope name F4

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 1/15 responders recognized this peptide, SI = 7.6.

HXB2 Location gp160 (324–336) Author Location Env (UG92005) Epitope GNIRQAHCNVSKA

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the V3 region of UG92005 (UG, clade D) and was recognized by two hybridoma with V $\beta$  usage V $\beta$ 8.2 and not determined.
- The epitope described here is the region of overlap of two 15 mers that were both able to stimulate IL-2 production from the hybridoma (IVGNIRQAHCNVSKA and GNIRQAHCN-VSKAKW)
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.

- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (324–338)

**Author Location** Env (UG92005)

Epitope GNIRQAHCNVSKAKW

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the V3 region of UG92005 (UG, clade D) and was recognized by eleven hybridomas with V $\beta$  usage V $\beta$ 5, 7, 8.1, 8.2, 11 and not determined a V $\beta$  8.1's and V $\beta$  8.2 also were shown to use V $\alpha$  8, and one of the ND used V $\alpha$  2.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and  $V\beta$  usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.

- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (324–338)

Author Location gp120 (V3)

Epitope GNIRQAHCNVSKAKW

Subtype B, D

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: Env Adjuvant: Complete Freund's

Adjuvant (CFA)

Species (MHC) mouse (H2b)

Assay type Cytokine production, CD4 T-cell Elispot -

IFNγ

**Keywords** vaccine-induced epitopes, variant cross-recognition or cross-neutralization, vaccine

antigen design **References** Zhan *et al.* 2004

- To develop a polyvalent Env vaccine, subtype specific B and D T-helper epitopes were identified, and mixtures of strain 1007, clade B, or UG92005, clade D envelopes were given to C57BL/6J mice. Mice were intramuscularly immunized with recombinant DNA, then intraperitoneally with rVV and finally with env protein in CFA. A dilution of 1/100 could still elicit type specific responses, and responses persisted with different prime boost combinations.
- Priming with 1007 and UG92005 env's induced both Envspecific (SNNTVGNPIILPCRI and AMYAPPIAGLIQCSS) and cross-reactive (PKVSFEPIPIHYCAP, C2, GNIRQAHCN-VSKAKW, V3-C3) Th responses in murine spleen cells.

**HXB2 Location** gp160 (327–341)

Author Location gp120 (327–341 HXB2)

Epitope RQAHCNISRAKWNNT

Subtype B

Immunogen vaccine

*Vector/Type:* protein *Strain:* B clade HXB2 *HIV component:* gp120

Species (MHC) mouse (I-A<sup>d</sup>)

**References** Warren & Thomas 1992

 Minimum epitope and MHC restriction determined for CTL clone that recognizes the N-terminal flank of the V3 loop.

**HXB2 Location** gp160 (327–346)

Author Location gp120 (331–350 MN)

Epitope RQAHCNISRAKWNDILRQIV

Epitope name 1988 Subtype B

Immunogen vaccine

Vector/Type: DNA, protein Strain: B clade MN HIV component: gp120 Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) guinea pig

**Keywords** vaccine-specific epitope characteristics, Th1 **References** Chattergoon *et al.* 2002

- Hartley guinea pigs were intradermally injected with either recombinant protein or plasmid expressed gp120 and monitored for delayed type hypersensivity (DTH) responses after vaccination, which are related to Th1 T-cell responses. CFA did not augment responses in animals vaccinated with plasmid.
- A total of 7 gp120 peptides elicited a delayed type hypersensivity (DTH) response after vaccination, out of a set of 60 overlapping peptides that spanned gp120. The vaccine delivery system, DNA versus rec protein, resulted in the recognition of distinct peptides.
- 4/5 animals vaccinated with rec gp120 responded by DTH to this peptide, and 2/6 responded that were vaccinated with plasmid gp120 DNA.

**HXB2 Location** gp160 (330–350) **Author Location** gp120 (330–349 IIIB)

Epitope HCNISRAKWNNTLKQIASKLR?

Epitope name F5 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 3/15 responders recognized this peptide, average SI = 5.5.

**HXB2 Location** gp160 (331–345) **Author Location** gp120 (IIIB)

Epitope CNISRAQWNNTLEQI

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro.
- Peptide priming does not always induce T-cells that recognize whole protein.

**HXB2 Location** gp160 (332–354) **Author Location** gp120 (337–359 NL43)

Epitope NISRAKWNATLKQIASKLREQFG

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 • HIV component: gp120, gp160

Species (MHC) human

References Sitz et al. 1999

• There was a great breadth of proliferative response to env peptides in 19 HIV-1 infected rgp160 and 17 HIV-1 infected rgp120 vaccine recipients.

• More than 30% of vaccinees had a stimulation index of greater than 5 to this peptide.

**HXB2 Location** gp160 (335–349) **Author Location** gp160 (342–356 IIIB)

Epitope RAKWNNTLKQIDSKL

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>b</sup>, H-2<sup>s</sup>)

**References** Berzofsky *et al.* 1991b; Berzofsky *et al.* 1991a • B10.BR (H-2A $^k$ , E $^k$ ), B10.A(5R) (H-2A $^b$ , E $^b$ ) and B10.S(9R)

(H-2A<sup>s</sup>, E<sup>s</sup>) mice immunized with rec gp160 showed a proliferative response to this peptide.

FVTIGKIGNMRQAHCNISRAKWNNTLKQIDSKL encompasses several murine Th epitopes including RAKWNNTLKQIDSKL and is referred to as a "multideterminant region" or cluster peptide.

**HXB2 Location** gp160 (335–349) **Author Location** gp120 (342–356 IIIB)

Epitope RAKWNNTLKQICSKL

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse  $(H-2^k, H-2^{t4}, H-2^{i5})$ 

References Hale et al. 1989

 Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

**HXB2 Location** gp160 (337–356) **Author Location** gp120 (341–360 MN)

Epitope KWNDTLRQIVSKLKEQFKNK

Epitope name 1989 Subtype B Immunogen vaccine

> Vector/Type: DNA, protein Strain: B clade MN HIV component: gp120 Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) guinea pig

**Keywords** vaccine-specific epitope characteristics, Th1 **References** Chattergoon *et al.* 2002

- Hartley guinea pig were intradermally injected with either recombinant protein or plasmid expressed gp120 and monitored
  for delayed type hypersensivity (DTH) responses after vaccination, which are related to Th1 T-cell responses. CFA did not
  augment responses in animals vaccinated with plasmid.
- A total of 7 gp120 peptides elicited a delayed type hypersensivity (DTH) response after vaccination, out of a set of 60 overlapping peptides that spanned gp120. The vaccine delivery system, DNA versus rec protein, resulted in the recognition of distinct peptides.
- 3/5 animals vaccinated with rec gp120 responded by DTH to this peptide, and 2/6 responded that were vaccinated with plasmid gp120 DNA.

**HXB2 Location** gp160 (339–359)

**Author Location** gp120 (340–359 89.6)

Epitope NNTLQQIVIKLREKFRNKTI

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse (H-2<sup>k</sup>, H-2<sup>d</sup>) Keywords immunodominance References Dai *et al.* 2001

- Promiscuous immunodominant epitopes in gp120 were mapped by overlapping peptides in CBA/J H-2<sup>k</sup> and BALB/c H-2<sup>d</sup> mice, and all were found to be in the outer domain, proximal to regions of structural disorder indicated by the crystal structure or by sequence divergence.
- This peptide was recognized by 4/10 CBA/J and 6/10 BALB/c mice with SI > 4, averaging 4.9 and 5.5 and is considered to be promiscuously immunodominant.
- Uniquely immunodominant sequences tended to be in the inner domain of the protein.

HXB2 Location gp160 (339–359)

Author Location gp120 (340–359 89.6)

Epitope NNTLQQIVIKLREKFRNKTI

Epitope name Peptide 32 Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse Donor MHC H-2k, H2-d

Keywords epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was reactive in 6/10 BALB/c mice tested, and in 4/10 CBA/J mice and was considered one of the 3 immunodominant peptides identified that were shared in both mouse strains: RRNIIGDIRQAHCNISRAKW, NNTLQQIVIKLREK-FRNKTI, GTNGTEGNDIITLQCRIKQI.

HXB2 Location gp160 (341–356) Author Location gp120 (IIIB)

Epitope TLEQIVKKLREQFGNC

**Immunogen** in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro
- Peptide priming does not always induce T-cells that recognize whole protein.

**HXB2 Location** gp160 (342–361)

Author Location gp120 (342–361 IIIB)

Epitope LKQIASKLREQFGNNKTIIF?

Epitope name G1 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 2/15 responders recognized this peptide, average SI = 6.0.

HXB2 Location gp160 (344–357)
Author Location gp120 (346–359)
Epitope QIVKKLREQFGNNK
Immunogen HIV-1 infection
Species (MHC) human

References Krowka et al. 1990

 Conjugation of HIV peptides to liposomes and rIL-2 stimulation may enhance cell-mediated responses.

HXB2 Location gp160 (349–368)

Author Location gp120 (350–369 89.6)

Epitope LREKFRNKTIAFNQSSGGD

**Epitope name** Peptide 33 **Immunogen** vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

**Species (MHC)** mouse **Donor MHC** H-2k, H2-d

Keywords epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was reactive in 3/10 BALB/c mice tested, and in 5/10 CBA/J mice.

**HXB2 Location** gp160 (350–370) **Author Location** gp120 (350–370 IIIB)

Epitope REQFGNNKTIIFKQSSGGDPE?

Epitope name G2

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

 Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.

- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 1/15 responders recognized this peptide, average SI = 3.2.

HXB2 Location gp160 (353-360) Author Location gp120 (355–362 IIIB)

Epitope FGNNKTII Immunogen SHIV infection

Species (MHC) macaque

References Lekutis & Letvin 1997

- C3 region minimal epitope determined through fine epitope mapping.
- Cell line was lost prior to confirmation of MHC requirements.

HXB2 Location gp160 (363–372) Author Location gp120 (368–377 LAI)

**Epitope** QSSGGDPEIV

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

HXB2 Location gp160 (364-378)

**Author Location** gp120 (364–378 IIIB, B10)

Epitope SSGGKPEIVTHSFNC

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (369–383)

**Author Location** gp120 (369–383 IIIB, B10)

**Epitope** PEIVTHSFNCGGEFF Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

monly evoke T-cell responses.

HXB2 Location gp160 (380-393) Author Location gp120 (380–393 IIIB)

Epitope GEFFYCNSTQLFNS?

Epitope name G4 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Keywords immunodominance

References Geretti et al. 1994

• Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.

- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- Five peptides were recognized most frequently: C2 (aa 142-161), C3 (aa 152-171), C5 (aa 172-191), E5 (aa 272-291) and G4 (aa 380-393). The first three were in or near V2, the other two were proximal to the V3 and V4 loops.
- 4/15 responders recognized this immunodominant peptide, average SI = 4.4.

HXB2 Location gp160 (381–395)

Author Location gp120 (IIIB)

**Epitope** EFFYCNTTQLFNNTW

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in
- Peptide priming does not always induce T-cells that recognize whole protein.

**HXB2 Location** gp160 (391–405)

Author Location gp120 (IIIB)

**Epitope** FNNTWRLNHTEGTKGC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (392-411)

Author Location gp120 (392–411 IIIB)

Epitope NSTWFNSTWSTEGSNNTEGS?

Epitope name G5

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- 12 gag and 18 env peptides were identified that could com-HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
  - · After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
  - IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
  - 1/15 responders recognized this peptide, SI = 9.3.

**HXB2 Location** gp160 (394–408)

**Author Location** gp120 (394–408 IIIB, B10)

**Epitope** TWFNSTWSTKGSNNT Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could com- Author Location Env (1007) monly evoke T-cell responses.

HXB2 Location gp160 (399-413)

**Author Location** gp120 (399–413 IIIB, B10)

Epitope TWSTKGSNNTEGSDT

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (404-423)

**Author Location** gp120 (400–419 89.6)

Epitope GTNGTEGNDIITLQCRIKQI

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup>) Keywords immunodominance References Dai et al. 2001

- Promiscuous immunodominant epitopes in gp120 were mapped by overlapping peptides in CBA/J H-2<sup>k</sup> and BALB/c H-2<sup>d</sup> mice, and all were found to be in the outer domain, proximal to regions of structural disorder indicated by the crystal structure or by sequence divergence.
- This peptide was recognized by 4/10 CBA/J and 6/10 BALB/c mice with SI > 4, averaging 4.9 and 5.5 and is considered to be promiscuously immunodominant.
- Uniquely immunodominant sequences tended to be in the inner domain of the protein.

HXB2 Location gp160 (404-423)

**Author Location** gp120 (400–419 89.6)

Epitope GTNGTEGNDIITLQCRIKQI

Epitope name Peptide 38 Immunogen vaccine

> Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse

Donor MHC H-2k, H2-d

Keywords epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was reactive in 8/10 BALB/c mice tested, and in 6/10 CBA/J mice, and was considered one of the 3 immunodominant peptides identified that were shared in both mouse strains: RRNIIGDIRQAHCNISRAKW, NNTLQQIVIKLREK-FRNKTI, GTNGTEGNDIITLQCRIKQI.

HXB2 Location gp160 (405–420)

Epitope SNNTVGNPIILPCRI

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete

Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

Keywords subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the V4C4 region of 1007 (US, clade B) and was recognized by 13 hybridomas with  $V\beta$  usage  $V\beta$ 4, 7, 8.1, 8.2, 10, 12 and not determined – one of the V $\beta$  8.2 was shown to utilize  $V\alpha$  2.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and  $V\beta$  usage was determined.
- · Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (405–420)

Author Location Env (gp160) (1007)

Epitope SNNTVGNPIILPCRI

Subtype B

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: Env Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (H2b)

Assay type Cytokine production, CD4 T-cell Elispot -

IFNγ

Keywords vaccine-induced epitopes, variant crossrecognition or cross-neutralization, vaccine

antigen design

References Zhan et al. 2004

- To develop a polyvalent Env vaccine, subtype specific B and D T-helper epitopes were identified, and mixtures of strain 1007, clade B, or UG92005, clade D envelopes were given to C57BL/6J mice. Mice were intramuscularly immunized with recombinant DNA, then intraperitoneally with rVV and finally with env protein in CFA. A dilution of 1/100 could still elicit type specific responses, and responses persisted with different prime boost combinations.
- T-cell hybridoma 1007P3-23 was isolated from mice immunized with 1007, and it recognized the peptide SNNTVGN-PIILPCRI of the V4/C4 region. The minimal, core peptide recognized by 10007P3-23 was NPIIL, a sequence not found in UG92005, which has a deletion in the core, so that the equivalent region in the D isolate is NNET—ITLQCRI
- Priming mixtures of 1007 and UG92005 induced both Envspecific (SNNTVGNPIILPCRI and AMYAPPIAGLIQCSS) and cross-reactive (PKVSFEPIPIHYCAP, C2, GNIRQAHCN-VSKAKW, V3-C3) Th responses in murine spleen cells.

HXB2 Location gp160 (410–429)

Author Location gp120 (410–429 PV22)

Epitope GSDTITLPCRIKQFINMWQE

Immunogen HIV-1 infection

**Species (MHC)** human (DR4) **References** Callahan *et al.* 1990

• Synthetic peptides representing natural variants were used to test for recognition in the context DR4.

HXB2 Location gp160 (410–429)
Author Location gp120 (410–429 PV22)
Epitope GSDTITLPCRIKQFINMWQE
Immunogen HIV-1 infection
Species (MHC) human (DR4(Dw10))
References Polydefkis et al. 1990

• Human CD4+ T-cell clones lyse recombinant vaccinia virusinfected cells that synthesize envelope gp160.

HXB2 Location gp160 (412–431)
Author Location gp120 (412–431 IIIB)
Epitope DTITLPCRIKQIINMWQKVG?
Epitope name H2
Subtype B
Immunogen HIV-1 infection

**Species** (MHC) human **References** Geretti *et al.* 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 1/15 responders recognized this peptide, SI = 5.7.

HXB2 Location gp160 (416–431)

Author Location gp120 (IIIB)

Epitope LPCRIKQIINMWQEVY

**Immunogen** in vitro stimulation or selection **Species (MHC)** human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro.
- Peptide priming does not always induce T-cells that recognize whole protein.

**HXB2 Location** gp160 (418–436) **Author Location** Env (417–435)

Epitope CRIKQIINMWQGVGKAMYA

Immunogen HIV-1 infection Species (MHC) human, chimpanzee References Nehete *et al.* 1998b

HIV-infected chimpanzees and HIV-positive patients show positive proliferative responses to multiple peptides from five conserved regions of the HIV-1 Env.

HXB2 Location gp160 (421–436)

Author Location gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection

Species (MHC) human (DR)
References Baier *et al.* 1995

 Linked HIV-1 T1 and P18 peptides to anti-HLA-DR and anti-IgD Fab fragments to enhance uptake by antigen presenting cells and thus increase immunogenicity.

**HXB2 Location** gp160 (421–436)

**Author Location** 

Epitope KQIINMWQEVGKAMYA

Epitope name T1
Immunogen vaccine

*Vector/Type:* canarypox prime with recombinant protein boost *HIV component:* gp120

Species (MHC) human (DRB1\*13)

Donor MHC DRB1\*01, DRB1\*13

Assay type proliferation

Keywords enhancing activity, optimal epitope

References Okazaki et al. 2006

- KQIINMWQEVGKAMYA-specific human CD4+ T-cell line from a healthy Caucasian American volunteer immunized with a canarypox virus vector expressing gp120 and boosted with recombinant gp120 was developed and found to be restricted to DR $\beta$ 1\*13. Epitope enhancement with different amino acid substitutions was studied.
- Likely binding core for KQIINMWQEVGKAMYA was determined as WQEVGKAMY, based on single A or S substitutions that diminished recognition, and proliferation assay data using truncated peptides.
- HLA binding motif was studied using substituted peptides in anchor positions 1,4,6,9 from the N-terminus of WQEVGKAMY. In position 1, CD4+ response was reduced by kqiinm[w/ai]QEVGKAMYa substitutions, but not by kqiinm[w/f]QEVGKAMYa substitution suggesting a requirement of aromatic amino acid. In position 4, CD4 response was reduced by kqiinmMWQE[v/af]GKAMYa substitutions, but

ing a requirement of aliphatic amino avid. In position 6, response was reduced by kqiinmWQEVG[k/ae]AMYa substitutions, but was enhanced by positively charged R substitution (kqiinmWQEVG[k/r]AMYa). In position 9, all peptides substituted with small, aromatic, or aliphatic amino acids (kqiinmWQEVGKAM[y/afi]a) induced enhanced response.

- The altered KOIINMWOE[v/i]GKAMYA peptide produced higher IFN-γ production that the original peptide, suggesting greater CD4 T-cell activation in a Th1 functional response.
- Triple substituted peptide KQIINMWQE[v/i]G[k/r]AM[y/a]A shifted the peak proliferative response to lower concentrations.

HXB2 Location gp160 (421–436) **Author Location** Env (421–436 IIIB) Epitope KQIINMWQEVGKAMYA

Epitope name T1 Immunogen vaccine

Vector/Type: peptide Strain: modified B clade IIIB HIV component: Env

Species (MHC) mouse (Ek)

Assay type Cytokine production, Th support of CTL response

Keywords binding affinity, Th1 References Ahlers et al. 2001

- BALB/c and A.AL were immunized with an Env-peptide vaccine construct containing the CTL epitope P18IIIB and a T helper epitope.
- Substitution of Glu (wt) to Ala, kqiinmwqAvgkamya, caused increased affinity for MHC class II Ek. This resulted in the upregulation of CD40L in the responding Th cells, and shifted the response towards Th1. Increased Th responses stimulated DCs to produce higher levels of IL-12, and B7-1 and B7-2, thus enhance CTL responses.
- The modified epitope, T1A, elicited stronger protection against increasing doses of viral challenge with vaccinia expressing HIV-1 IIIB gp120 compared to the wildtype epitope T1.

HXB2 Location gp160 (421-436) Author Location gp120 (428-443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1 Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Klinman et al. 1995

• Hybrid T1-V3 peptide activates IL-4 and IL-6 in a dose dependent manner.

HXB2 Location gp160 (421-436) Author Location gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1 Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

**Species (MHC)** mouse (H-2<sup>k</sup>) References Ahlers et al. 1997b

• first identified Th epitope in HIV.

- enhanced by kqiinmWOE[v/i]GKAMYa substitution, suggest• Alanine at position 436 (instead of E in wild-type) enhances MHC binding and antigenicity of peptide by several orders of magnitude.
  - Vaccines with a CTL epitope linked to a more potent helper epitope yielded greatly enhanced CTL response relative to the wildtype helper epitope.
  - T1 peptide linked to CTL epitopes in four vaccine constructs used to immunize mice: KOIIN-MWQEVGKAMYAPPISGQIRRIQRGPGRAFVTIGK, KQIINMWQEVGKAMYAPPISGQIRRIQRGPGRAFVTI, KQIINMWQAVGKAMYAPPISGQIRRIQRGPGRAFVTIGK, KQIINMWQAVGKAMYAPPISGQIRRIQRGPGRAFVTI.

HXB2 Location gp160 (421-436)

**Author Location** gp120 (428–443 IIIB, B10)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen computer prediction

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup>, H-2<sup>s</sup>)

References Cease et al. 1987

• 1 of 2 functional epitopes identified using an amphipathic helix epitope prediction algorithm.

**HXB2 Location** gp160 (421–436) Author Location gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1 Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup>, H-2<sup>t4</sup>)

References Hale et al. 1989

• Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

HXB2 Location gp160 (421-436)

Author Location gp160 (428-443 IIIB)

Epitope KQIINMWQEVGKAMYA

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (H-2<sup>k</sup>, H-2<sup>s</sup>, H-2<sup>d</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- B10.BR (H-2 $A^k$ ,  $E^k$ ), B10.D2 (H-2 $A^d$ ,  $E^d$ ) and B10.S(9R) (H-2As, Es) mice immunized with rec gp160 showed a proliferative response to this peptide.
- KQIINMWQEVGKAMYAPPISGQIR encompasses several murine Th epitopes including KQIINMWQEVGKAMYA and is referred to as a "multideterminant region" or cluster peptide.

**HXB2 Location** gp160 (421–436)

Author Location gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1 Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

**Species** (MHC) mouse (H-2E $\alpha$  E $\beta$  <sup>k</sup>)

References Boehncke et al. 1993

- C3H H2<sup>k</sup> mice were used for immunization in the study because H-2<sup>k</sup> mice are particularly good T1 responders T1 can be presented by  $E\alpha E\beta^k$  but not  $E\alpha E\beta^b$  the nature of the T1 class II molecular interaction was thoroughly explored.
- Alanine substitutions across peptide did not negatively affect MHC binding or effective presentation of epitope, except at three critical residues (432N, 435Q, 439K), however substitutions with larger side chains often diminished activity – only a few amino acids were found to be critical for class II interaction and for maintaining T-cell receptor specificity.
- A gain in potency was observed when 436E was replaced with A, suggesting that substitutions in positions that interfere with binding might allow the design of a more potent vaccine.

HXB2 Location gp160 (421–436) Author Location gp120 (426–441 IIIB) Epitope KQFINMWQEWGKAMYA

Immunogen

Species (MHC) human

References Furci et al. 1997

- Epitope T1 variant: 9/11 exposed-uninfected individuals in this study had a proliferative response to a C5 peptide, but none reacted with this previously defined epitope.
- IIIB position 435 listed as W in this epitope as opposed to V in the sequence.

**HXB2 Location** gp160 (421–436) **Author Location** gp120 (428–433 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection

Species (MHC) human

Keywords responses in children, kinetics, Th1

References Wasik et al. 2000

- Th responses measured by IL-2 responses to P18 and T1 in HIV-1 infected infants were undetectable at less than 1 month of age, and remained low in children with AIDS symptoms, but increased with age in children with slowly progressive disease.
- The kinetics and intensity of the CTL activity during the first year of life was related to the child's ability to make Th1 responses.

**HXB2 Location** gp160 (421–436) **Author Location** gp120 (428–433 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection

Species (MHC) human

Keywords responses in children, Th1, Th2

References Wasik et al. 1997

- The breadth and intensity of the CTL response and the type of Th response was studied in seven rapidly progressing HIV-1 + infants.
- IL-2 and γ IFN production from Th1 cells correlated with the CTLp frequency against HIV-1 Gag, Env, Nef and Pol.
- IL-4 production from Th2 cells was inversely correlated with the CTLp frequency.
- The HIV-1 + children with strong CTL responses had levels of anti-CD3 MAb induction of Th1 cells comparable to those of uninfected children.

**HXB2 Location** gp160 (421–436)

Author Location gp120 (428-443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1
Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

Species (MHC) human

References Berzofsky et al. 1988

 Proliferative response to T1 and T2 peptides in 14 immunized, uninfected humans.

HXB2 Location gp160 (421–436)

Author Location gp120 (428-443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

Species (MHC) goat

References Palker et al. 1989

 Goats immunized with peptides containing V3 type-specific neutralizing determinants coupled to T1.

**HXB2 Location** gp160 (421–436)

Author Location gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1989

IL-2 production detection of Th lymphocytes from asymptomatic HIV-positive individuals.

**HXB2 Location** gp160 (421–436)

Author Location gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1991a

 Peptides stimulate Th cell function and CTL activity in similar patient populations.

HXB2 Location gp160 (421-436)

**Author Location** gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp160

Species (MHC) human

References Clerici et al. 1991b

• Immunizing uninfected individuals with rgp160 results in stronger Th response than does natural infection.

**HXB2 Location** gp160 (421–436)

Author Location gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

**Immunogen** 

Species (MHC) human

References Clerici et al. 1992

 Cell-mediated immune response to HIV-1 peptides in HIV-1 exposed seronegative men.

HXB2 Location gp160 (421–436) Author Location gp120 (428–443 IIIB) Epitope KQIINMWQEVGKAMYA

Immunogen vaccine

Vector/Type: bacteriophage coat protein Strain: B clade MN HIV component: V3

Species (MHC) mouse

References di Marzo Veronese et al. 1994

• Epitope T1 was engineered into a filamentous bacteriophage coat protein, and the Th epitope stimulated Ab production to the V3 loop.

HXB2 Location gp160 (421–436) Author Location gp120 (428–443 IIIB) Epitope KQIINMWQEVGKAMYA

Epitope name T1
Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

**Species** (MHC) chimpanzee **References** Haynes *et al.* 1993

 Hybrid T1-V3 peptide immunogenicity reduced when the fusogenic domain of gp41 was added.

HXB2 Location gp160 (421–436) Author Location gp120 (428–443 IIIB) Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1997

 Used in a study of the influence of pentoxifylline on HIV specific T-cells.

HXB2 Location gp160 (421–436) Author Location gp120 (428–443 IIIB) Epitope KQIINMWQEVGKAMYA

Epitope name T1 Immunogen Species (MHC) human

References Pinto et al. 1995

 CTL activity analyzed in parallel with Th reactivity in exposed but uninfected health care workers.

HXB2 Location gp160 (421–436) Author Location gp160 (428–433 IIIB) Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human Keywords immunodominance

References Wasik et al. 1999

• IL-2 responses associated with beta-chemokine expression were detectable at birth in the majority of uninfected infants born to HIV+ mothers, declining by age 6 months.

- T1 peptide: In both uninfected and infected infants of HIVpositive mothers, responses to the T1 peptide were more frequent than responses to P18 (RIQRGPGRAFVTIGK)
- T1 is a highly conserved epitope, whereas P18 has a higher mutation rate due to its location in the immunodominant V3 loop region.

**HXB2 Location** gp160 (421–436) **Author Location** gp120 (428–443 IIIB)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection

Species (MHC) human

References Kaul et al. 1999

- Kenyan sex workers that remained seronegative were found to frequently have HIV-env peptide specific Th responses detected by an IL-2 assay (11/20 cases) and mucosal genital tract anti-HIV IgA (16/21 cases)
- Helper epitopes used in this study were noted to be previously described Clerici et al. [1989], and were not explicitly described in Kaul et al. [1999]

HXB2 Location gp160 (421–436) Author Location gp120 (MN)

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide Strain: B clade MN

Species (MHC) human

References Bartlett et al. 1998

- C4-V3 PV (polyvalent HIV envelope synthetic peptide immunogen) consisted of T1 helper epitope presented in tandem with a V3 loop CTL epitope from one of four different North American strains.
- This was a pilot phase I study involving vaccination of ten HIV-infected subjects who were HLA-B7-positive.
- Enhanced lymphoproliferative response to peptide was observed in 5/8 vaccinees increase in neutralizing antibody responses in 4/8 vacinees.

HXB2 Location gp160 (421-436)

Author Location gp120

Epitope KQIINMWQEVGKAMYA

Epitope name T1

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human

**Keywords** subtype comparisons, responses in children, mother-to-infant transmission

References Kuhn et al. 2001a

- In a S. African perinatal transmission study, 33% (33/86) of cord blood samples from infants with seropositive mothers produced T-helper responses (measured by a bioassay measuring IL2 production in a murine cell line and confirmed with a proliferation assay) against a peptide cocktail containing The epitopes P18 MN, P18 IIIB, T1, T2, and TH4.
- The mothers were predominantly infected with subtype C, but the T help response was detectable in a number of cord blood samples despite using peptides based on B subtype reagents.

- 3/33 infants with cord blood T help responses to Env were infected in utero, 2/33 were lost to follow up, and 28/33 were not infected. 6/53 of the infants with cord blood that was unresponsive to Env peptide stimulation were infected before delivery, and 8/47 contracted HIV intrapartum or via breastfeeding.
- Measurable HIV specific T help responses elicited in the immunologically immature newborn, possibly in response to in utero exposure, are associated with a protective natural immunity that helps block mother-infant transmission of HIV-1.

HXB2 Location gp160 (421-436) Author Location gp120 (428-443 RF) Epitope KQIINMWQEVGKAMYA Epitope name T1

Immunogen HIV-1 infection

Species (MHC)

Keywords epitope processing References de Lorimier et al. 1994

- Proton NMR spectroscopy was employed to analyze the solution conformation of a hybrid peptide, T1-SP10RF, in order to better understand the immunogenicity of its' T helper (KQI-INMWQEVGKAMYA, CTRPNNNTRKSI), CTL (SITKGP-GRVIYATG) and B-cell epitopes (RKSITKGPGRVIYATG).
- As a free peptide, the T1 segment, a T-helper epitope is in an extended conformation with nascent helical conformation. It may form a beta strand in native gp120, and a nonnative conformation may account for the inability of free T1 peptide to elicit antibody responses, in contrast to the T1 segment in native gp120. It lacks random-coil conformations, and it is suggested that this may make the peptide less susceptible to complete proteolytic degredation, and be favored within epitopes.

HXB2 Location gp160 (421-436) Author Location Env (428–443 IIIB) Epitope KQIINMWQEVGKAMYA

Epitope name T1

Species (MHC)

Assay type Cytokine production

Keywords mother-to-infant transmission

References Clerici et al. 1993a

- Cord blood samples in 8/23 infants with HIV+ mothers showed IL-2 production in response to peptides from HIV-1 gp1260, demonstrating that Th cell priming to HIV env determinants can occur in utero. HIV-specific Th immunity in utero may be protective, as none of the 8 with HIV-1 specific Th activity became infected, while 3/15 infants with no detectable Th activited were infected.
- PBL from 10/21 of the mothers showed HIV-1 specific Th activity through IL-2 production in response to Env peptides.

HXB2 Location gp160 (421-436) **Author Location** Env (IIIB) Epitope KQIINMWQEVGKAMYA Epitope name T1 Subtype B Immunogen HIV-1 exposed seronegative Species (MHC)

Assay type Cytokine production References Clerici et al. 1994a

- Six of eight HIV-exposed health care workers had transient HIV-specific T-helper responses after percutaneous exposure to HIV, responding to two or more Env peptides among the set T1, T2, Th4.1, P18IIIB and P18MN. Responses were lost after 12-56 weeks. The HIV-specific Th responses occurred without seroconversion or PCR evidence for infection.
- Six of the eight HIV-exposed individuals responded to two or more peptides, whereas, only one individual exposed to HIV-negative blood responded to one peptide.

HXB2 Location gp160 (421–436) **Author Location** HIV-1 (IIIB) Epitope KQIINMWQEVGKAMYA

Epitope name T1 Subtype B

Immunogen HIV-1 infection

Species (MHC)

Assay type Cytokine production References Clerici et al. 1994b

• IL-10-specific mRNA was upregulated in PBMC from asymptomatic, HIV-infected (HIV+) patients, and was particularly high in those with severely compromised Th cells function. Th response to HIV peptides in vitro could be restored by IL-10

HXB2 Location gp160 (421–436) **Author Location** Env (428–443)

Epitope KQIINMWQEVGKAMYA

Epitope name T1 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type proliferation

**Keywords** responses in children, mother-to-infant transmission

References Kuhn et al. 2001b

- **Immunogen** HIV-1 infection, HIV-1 exposed seronegative T helper proliferative responses in cord blood at delivery to a cocktail of HIV envelope peptides were infrequent (1/41) among infants whose HIV-positive mothers received antiretroviral prophylaxis at delivery to prevent transmission. Responses were 10 times more frequent among infants of HIV-seropositive women who had no antiretroviral treatment (7/29). Reductions of HIV-specific responses occurred despite persistance of detectable HIV RNA in the mothers at delivery.
  - The reduction of Th responses in newborns raises the possibility that anti-retrovial exposure during pregnancy may block subsequent immune protection. The authors point two relevant citations that showed ARV exposed infants had a more rapid course of progression (Kuhn et al., JID 182:104 (2000)), but were at no greater risk of infection due to subsequent breast feeding (Ditrame et al., Lancet 354:2050 (1999)).

HXB2 Location gp160 (421–436) **Author Location** Env (gp160) (421–436) Epitope KQIINMWQEVGKAMYA Epitope name T1 Subtype C Immunogen HIV-1 infection

Species (MHC) human

Country South Africa.

Assay type proliferation

**Keywords** responses in children, variant crossrecognition or cross-neutralization

References Meddows-Taylor et al. 2004

- Viral isolates (gp160) from 16 vertically HIV-1 infected children (8 T-helper cell Env responders and 8 non-responders) were analyzed for variation after comparison with Env peptide sequences (conserved regions T1, T2 and TH4.1, hypervariable regions P18 MN and P181 IIIB) used for in vitro stimulation.
- · No correlation between the age, clinical category (mild or severe), HIV-1 viral load and the degree of epitope variation was established.

HXB2 Location gp160 (421-444)

Author Location gp160 (428–451 IIIB)

Epitope KQIINMWQEVGKAMYAPPISGQIR

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** human, mouse (H-2<sup>k</sup>, H-2<sup>b</sup>, H-2<sup>s</sup>, H-2<sup>d</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- KQIINMWQEVGKAMYAPPISGQIR encompasses several murine Th epitopes and is referred to as a "multideterminant region" or cluster peptide.
- Six multideterminant region cluster peptides were evaluated Th responses in different MHC/HLA backgrounds after vaccination of mice with gp160, or in infected people.
- This cluster peptide elicited proliferative responses in cells from all H-2 haplotypes tested: B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>), B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), B10.A(5R) mice (H-2A<sup>b</sup>, E<sup>b</sup>), and B10.S(9R) mice (H-2A<sup>s</sup>, E<sup>s</sup>)
- IL-2 production in response to this peptide was observed in 73% (8/11) of asymptomatic HIV-infected individuals.

HXB2 Location gp160 (421-444)

Author Location gp120 (428–451 IIIB)

Epitope KQIIMNWQEVGKAMYAPPISGQIR

Epitope name T1

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

**Species (MHC)** mouse (H2<sup>d</sup>)

References Shirai et al. 1996a

• Linked to a CTL epitope from hepatitis C virus, induced CD4+ helper cells producing IL-2.

**HXB2 Location** gp160 (421–444)

**Author Location** Env (gp160) (HIV-1 IIIB)

Epitope KQIINMWQEVGKAMYAPPISGQIR

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB HIV component: Env Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72), Montanide (ISA 51)

Species (MHC) macaque

Assay type proliferation Keywords mucosal immunity References Belyakov et al. 2001

- Intrarectal vaccination with a Th and CTL peptide vaccine provided better protection against intrarectal challenge with pathogenic SHIV-Ku1 than subcutaneous administered vaccine. In some animals after the initial viremia, viral loads were diminished to undetectable levels in the blood and intestine, and CD4+ T cells were better preserved.
- The CD4 T-cell proliferative response correlated with the level of the CTL response.

HXB2 Location gp160 (423-440)

**Author Location** gp120 (428–445)

Epitope FINMWQEVGKAMYAPPIS

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression References Caruso et al. 1997

- · As HIV-1-infected individuals progress to disease, T-cells show reduced ability to proliferate in response to HIV antigen, but retain the ability to express the activation antigens CD25 and CD71.
- The ability to express activation markers in response to HIV is retained, but the response to tetanus toxoid recall antigen is
- This study investigated CD25 and CD71 expression in PBMC from patients at various stages of progression, measuring the response to in vitro stimulation by peptide cocktail containing four antigenic Env peptides, or p17 and p24.

**HXB2 Location** gp160 (424–438)

**Author Location** gp120 (424–438 IIIB, B10)

Epitope INMWQEVGKAMYAPP

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (425-439)

Author Location gp160 (432–446 IIIB)

Epitope NMWQEVGKAMYAPPI

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (H-2s)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- B10.S(9R) (H-2A<sup>s</sup>, E<sup>s</sup>) mice immunized with rec gp160 showed a proliferative response to this peptide.
- KQIINMWQEVGKAMYAPPISGQIR encompasses several murine Th epitopes including NMWQEVGKAMYAPPI and is referred to as a "multideterminant region" or cluster peptide.

**HXB2 Location** gp160 (425–439)

**Author Location** gp120 (432–446 IIIB)

Epitope NMWQEVGKAMYAPPI

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>t4</sup>) References Hale et al. 1989

 Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

HXB2 Location gp160 (426–441)

Author Location gp120 (IIIB)

Enitone MM/DEVGKAMYAPP

Epitope MWQEVGKAMYAPPIGC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro.
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (430–444) Author Location gp160 (437–451 IIIB) Epitope VGKAMYAPPISGQIR

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>b</sup>, H-2<sup>s</sup>, H-2<sup>d</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- This peptide elicited proliferative responses in cells from all H-2 haplotypes tested: B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>), B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), B10.A(5R) mice (H-2A<sup>b</sup>, E<sup>b</sup>), and B10.S(9R) mice (H-2A<sup>s</sup>, E<sup>s</sup>)
- KQIINMWQEVGKAMYAPPISGQIR encompasses several murine Th epitopes including VGKAMYAPPISGQIR and is referred to as a "multideterminant region" or cluster peptide.

HXB2 Location gp160 (430–444) Author Location gp120 (437–451 IIIB) Epitope VGKAMYAPPISGQIR

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup> H-2<sup>i5</sup>, H-2<sup>t4</sup>)

References Hale et al. 1989

• Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

**HXB2 Location** gp160 (430–453) **Author Location** gp120 (430–453)

Epitope VGKAMYAPPISGQIRCSSNITGLL

Immunogen vaccine

Vector/Type: protein HIV component: gp160

**Species (MHC)** mouse (H-2<sup>b</sup>)

**Keywords** epitope processing **References** Sjolander *et al.* 1996

- Study demonstrates that T-cell determinants from glycoproteins can depend on the glycosylation of the protein.
- Peptide stimulation of an *in vitro* proliferative response required *in vivo* priming with glycosylated protein.
- Local glycosylation sites thought not to be part of the epitope, but may be important for epitope processing.

HXB2 Location gp160 (432–451)

Author Location gp120 (432–451 IIIB)

Epitope KAMYAPPISGQIRCSSNITG?

Epitope name H4 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 1/15 responders recognized this peptide, SI = 6.3.

HXB2 Location gp160 (433–447) Author Location Env (UG92005) Epitope AMYAPPIAGLIQCSS

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This epitope is located in the C4 region of UG92005 (UG, clade D) and was recognized by ten hybridomas with  $V\beta$  usage  $V\beta$  6, 8.1, 8.2, 13, 14 and not determined among the ND  $V\beta$  set, three  $V\alpha$ s were identified,  $V\alpha$  2, 8, and 11.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site

through processing pathways.

**HXB2 Location** gp160 (433–447) Author Location Env (gp160) (UG92005) Epitope AMYAPPIAGLIQCSS

> Subtype D Immunogen vaccine

> > Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: Env Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (H2b)

Assay type Cytokine production, CD4 T-cell Elispot -IFNγ

Keywords vaccine-induced epitopes, variant crossrecognition or cross-neutralization, vaccine antigen design

References Zhan et al. 2004

- To develop a polyvalent Env vaccine, subtype specific B and D T-helper epitopes were identified, and mixtures of strain 1007, clade B, or UG92005, clade D envelopes were given to C57BL/6J mice. Mice were intramuscularly immunized with recombinant DNA, then intraperitoneally with rVV and finally with env protein in CFA. A dilution of 1/100 could still elicit type specific responses, and responses persisted with different prime boost combinations.
- T-cell hybridoma UGP2-17 was isolated from mice immunized with env sequence UG92005 (clade D), and it recognized the C4/V4 region peptide AMYAPPIAGLIQCSS. The minimal peptide recognized by 10007P3-23 was PPIAGLIQ, which matched only 5/8 residues in the B clade isolate, ppiRgQiK.
- Priming with 1007 and UG env's induced both Env-specific (SNNTVGNPIILPCRI and AMYAPPIAGLIQCSS) and crossreactive (PKVSFEPIPIHYCAP, C2, GNIRQAHCNVSKAKW, V3-C3) Th responses in murine spleen cells.

HXB2 Location gp160 (436-451) Author Location gp120 (IIIB)

**Epitope** APPIGGQISCSSNITY

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (438–460) **Author Location** gp120 (443–465 NL43)

Epitope PISGQIRCSSNITGLLLTRDGGN

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: gp120, gp160

Species (MHC) human

References Sitz et al. 1999

• There was a great breadth of proliferative response to env peptides in 19 HIV-1 infected rgp160 and 17 HIV-1 infected rgp120 vaccine recipients.

proximity may allow binding to lectins and promote trafficking • Close to 40% of vaccinees had a stimulation index of greater than 5 to this peptide.

HXB2 Location gp160 (439-448)

Author Location gp120 (151-160 W6.ID)

Epitope IGGQIRCSSN Immunogen vaccine

> Vector/Type: protein Strain: B clade W61D HIV component: gp120 Adjuvant: MPL-SE

adjuvant, QS21

Species (MHC) human

References Jones et al. 1999

- HIV-1 specific T-cell lines isolated from an HIV seronegative volunteer vaccinated with rgp120 and a QS21/MPL adjuvant.
- One T-cell line responds to two overlapping peptides, and the region of overlap is IGGQIRCSSN.
- The IIIB version of the first reactive peptide, EVGKAMYAP-PIGGQIRCSSN, has a single substitution and induces proliferation as well as the original W61D peptide: evgkamyappiSggircssn.

**HXB2 Location** gp160 (439–461)

**Author Location** Env (438–460)

Epitope IRGQIRCSSNITGLLLTRDGGNN

Epitope name HIV\_env\_DRB0101\_1

Subtype M

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type T-cell Elispot

**Keywords** computational epitope prediction

References De Groot et al. 2004

- Bioinformatic tools (EpiMatrix and Conservatrix) were employed to predict 9 highly conserved and promiscuous class II restricted T helper epitopes from HIV-1 env sequences for a cross-clade polyepitope vaccine. 10/34 HIV-1 infected patients induced Th1 immune responses to 8/9 study peptides.
- 2/28 subjects tested responded to this peptide.
- Class II peptides containing putative epitopes were selected to favor DRB0101 presentation, as there is a transgenic mouse model for this allele for future vaccine studies. The highest scoring DRB0101 sequence within this peptide was QIRCSS-NIT.

HXB2 Location gp160 (446–461)

Author Location gp120 (IIIB)

**Epitope** SSNITGLLLTRDGGTC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (452–471)

Author Location gp120 (452-471 IIIB)

**Epitope** LLLTRDGGNSNNESEIFRPG?

Epitope name 11 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 2/15 responders recognized this peptide, average SI = 3.5.

**HXB2 Location** gp160 (456–470)

Author Location gp120 (IIIB)

Epitope RDGGTNVTNDTEVFRC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro.
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (459-473)

**Author Location** gp120 (459–473 IIIB, B10)

 ${\bf Epitope} \ {\tt GNSNNESEIFRPGGG}$ 

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (468–483)

**Author Location** gp120 (466-481)

Epitope FRPGGGDMRDNWRSEL

Immunogen HIV-1 infection

Species (MHC) human

References Krowka et al. 1990

• Conjugation of HIV peptides to liposomes and rIL-2 stimulation may enhance cell-mediated responses.

**HXB2 Location** gp160 (472–491)

**Author Location** gp120 (472–491 IIIB)

Epitope GGDMRDNWRSELYKYKVVKI?

Epitope name I3

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.

- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 2/15 responders recognized this peptide, average SI = 7.2.

HXB2 Location gp160 (474–488)

**Author Location** gp120 (474–488 IIIB, B10)

Epitope DMRDNWRSELYKYKV

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (476–490)

Author Location gp120 (483-497 IIIB)

Epitope RDNWRSELYKYKVVK

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species** (MHC) mouse (H-2<sup>d</sup>, H-2<sup>t4</sup>)

References Hale et al. 1989

• Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

**HXB2 Location** gp160 (476–490)

Author Location gp160 (483-497 IIIB)

Epitope RDNWRSELYKYKVVK

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Com-

plete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>s</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- • This peptide elicited proliferative responses in B10.BR mice  $(H-2A^k \text{ and B10.S(9R) mice } (H-2A^s, E^s)$
- RDNWRSELYKYKVVKIEPLGVAPT encompasses several murine Th epitopes including RDNWRSELYKYKVVK and is referred to as a "multideterminant region" or cluster peptide.

**HXB2 Location** gp160 (476–499)

**Author Location** gp160 (483–506 IIIB)

Epitope RDNWRSELYKYKVVKIEPLGVAPT

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Com-

plete Freund's Adjuvant (CFA)

**Species (MHC)** human, mouse (H-2<sup>k</sup>, H-2<sup>b</sup>, H-2<sup>s</sup>, H-2<sup>d</sup>)

**References** Berzofsky et al. 1991b; Berzofsky et al. 1991a

- RDNWRSELYKYKVVKIEPLGVAPT encompasses several murine Th epitopes and is referred to as a "multideterminant region" or cluster peptide.
- Six multideterminant region cluster peptides were evaluated Th responses in different MHC/HLA backgrounds after vaccination of mice with gp160, or in infected people.
- This cluster peptide elicited proliferative responses in cells from all H-2 haplotypes tested: B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>), B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), B10.A(5R) mice (H-2A<sup>b</sup>, E<sup>b</sup>), and B10.S(9R) mice (H-2A<sup>s</sup>, E<sup>s</sup>)
- IL-2 production in response to this peptide was observed in 52% (14/27) of asymptomatic HIV-infected individuals.

HXB2 Location gp160 (479-498) Author Location gp120 (481–500 MN) Epitope WRSELYKYKVVTIEPLGVAP

Epitope name 2013 Subtype B Immunogen vaccine

> Vector/Type: DNA, protein Strain: B clade MN HIV component: gp120 Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) guinea pig

**Keywords** vaccine-specific epitope characteristics, Th1 References Chattergoon et al. 2002

- Hartley guinea pigs were intradermally injected with either recombinant protein or plasmid expressed gp120 and monitored for delayed type hypersensivity (DTH) responses after vaccination, which are related to Th1 T-cell responses. CFA did not augment responses in animals vaccinated with plasmid.
- A total of 7 gp120 peptides elicited a delayed type hypersensivity (DTH) response after vaccination, out of a set of 60 overlapping peptides that spanned gp120. The vaccine delivery system, DNA versus rec protein, resulted in the recognition of distinct peptides.
- 0/5 animals vaccinated with rec gp120 responded by DTH to this peptide, while 6/6 vaccinated with plasmid gp120 DNA responded.

HXB2 Location gp160 (482-501) Author Location gp120 (482–501 IIIB) Epitope ELYKYKVVKIEPLGVAPTKA

Immunogen vaccine

HIV component: Env

Species (MHC) macaque

References Lekutis et al. 1997

- tope in a rhesus monkey.
- Epitope was recognized by both monkeys used in this study.

HXB2 Location gp160 (482-501) Author Location gp120 (482–501 IIIB) Epitope ELYKYKVVKIEPLGVAPTKA?

Epitope name I4 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 3/15 responders recognized this peptide, average SI = 6.0.

HXB2 Location gp160 (483-502) **Author Location** gp120 (480–499 89.6) Epitope LYKYKVVRIEPIGVAPTRAK

Epitope name Peptide 46 Immunogen vaccine

> Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse Donor MHC H-2k, H2-d

> Keywords epitope processing, immunodominance References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was reactive in 7/10 BALB/c mice tested, and in only 1/10 CBA/J mice.

HXB2 Location gp160 (484–496) Author Location gp120 (484-496 HXB2)

Epitope YKYKVVKIEPLGV

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade HXB2

HIV component: Env

Species (MHC) macaque (DR\*W201) References Lekutis & Letvin 1998

- Vector/Type: DNA Strain: B clade IIIB Variants of this epitope with substitutions at position 490(K) retained ability to bind to MHC class II, but failed to induce proliferation/cytokine secretion in HIV-1 env-specific CD4+ Th cells.
- HIV-1 env DNA vaccine induced Th cell response to this epiinduced proliferative response.

HXB2 Location gp160 (484–498)

**Author Location** gp120 (484–498 IIIB, B10)

Epitope YKYKVVKIEPLGVAP Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (484–499)

Author Location gp120 (492–506 IIIB)

Epitope CKYKVVKIEPLGVAPT

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>k</sup>, H-2<sup>t4</sup>, H-2<sup>i5</sup>)

References Hale et al. 1989

· Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

**HXB2 Location** gp160 (485–499) Author Location gp160 (492-506 IIIB) **Epitope** KYKVVKIEPLGVAPT

Immunogen vaccine

HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>b</sup>, H-2<sup>s</sup>, H-2<sup>d</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- This peptide elicited proliferative responses in cells from all H-2 haplotypes tested: B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>), B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), B10.A(5R) mice (H-2A<sup>b</sup>, E<sup>b</sup>), and B10.S(9R) mice (H-2A<sup>s</sup>, E<sup>s</sup>)
- RDNWRSELYKYKVVKIEPLGVAPT encompasses several murine Th epitopes including KYKVVKIEPLGVAPT and is referred to as a "multideterminant region" or cluster peptide.

HXB2 Location gp160 (485-500) Author Location gp120 (IIIB)

Epitope KYKVIKIEPLGIAPTC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro.
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (486-494) Author Location gp120 (486–494 IIIB) **Epitope** YKVVKIEPL Immunogen SHIV infection

Species (MHC) macaque (DRB\*W201) **References** Lekutis & Letvin 1997

• C5 region minimal epitope determined through fine epitope mapping.

**HXB2 Location** gp160 (487–512) Author Location gp120 (494–518 IIIB)

Epitope KVVKIEPLGVAPTKAKRRVVQREKRC

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

Species (MHC) mouse

References Goodman-Snitkoff et al. 1990

• Identification of putative Th epitopes that stimulate an antibody response in peptide immunized mice.

HXB2 Location gp160 (492-512) Author Location gp120 (492-512 IIIB)

Epitope EPLGVAPTKAKRRVVQREKRA?

Epitope name 15 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- · After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.

- Vector/Type: protein Strain: B clade IIIB IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
  - 1/15 responders recognized this peptide, SI = 4.9.

HXB2 Location gp160 (493-511) **Author Location** gp120 (490–508 89.6) Epitope PIGVAPTRAKRRTVQREKR

Epitope name Peptide 47 Immunogen vaccine

> Vector/Type: protein Strain: B clade 89.6 HIV component: gp120 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (MHC) mouse

Donor MHC H-2k, H2-d

Keywords epitope processing, immunodominance

References Dai et al. 2001

- Helper T-cell proliferative responses to gp120 vaccines in 2 mouse strains, CBA/J and BALB/c, were mapped using 47 overlapping peptides that span gp120. Promiscuously immunodominant peptides were identified in both mice strains that were located in the outer domain of gp120 and were associated with regions of local structural disorder in proximal N-terminal segments, suggesting 3-D protein structure influences Th antigen processing and the frequency of immunogenic sequences.
- This peptide was reactive in only 2/10 BALB/c mice tested, and in 8/10 CBA/J mice.

HXB2 Location gp160 (499-511)

Author Location gp120 (IIIB)

**Epitope** TKAKRRVVEREKR

Immunogen in vitro stimulation or selection

Species (MHC) human (DR) References Wilson et al. 1997b

- Thought to be a mimic of a HLA class II DR  $\beta$  chain variable region.
- · Response to this epitope may cause a breakdown of selftolerance.
- Presentation of epitope induced autoreactive T-cell lines in PBMC from uninfected donors.
- Suppression of proliferation to soluble antigens by the CD8+ fraction of TKAKRRVVEREKR stimulated T-cells was observed.

HXB2 Location gp160 (499–519)

Author Location gp41 (MN)

Epitope TKAKRRVVQREKRAAIGALF

Epitope name TF20 Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type Cytokine production, proliferation, CD4 Tcell Elispot - IFN $\gamma$ , Intracellular cytokine

staining

Keywords HAART, ART, acute/early infection

References Malhotra et al. 2003

• 92 acute- or early-HIV infected subjects were tested for Envand Gag-specific Th responses. There was an overall low probability of detecting HIV-1-specific Th responses, Env responses were rare and (5%) and only detected very early in infection, probably reflecting a low frequency of response and impaired lymophoproliferative capacity, not viral escape. Gagspecific Th responses were observed more frequently (16%), were found throughout the acute and early infection phases, and predominated after ARV therapy.

 This epitope is one of six Env-specific Th responses detected in one patient 11 days post-infection that were studied in detail.
 These Th reponses were not detected at subsequent time points, despite the Env specific Th cells being maintained as detected by TCR clonal transcripts, and the epitope sequence being maintained in Env.

**HXB2 Location** gp160 (519–543) **Author Location** gp41 (519–543)

Epitope FLGFLGAAGSTMGAASLTLTVQARC

Immunogen vaccine

Vector/Type: peptide

**Species** (MHC) mouse (H-2<sup>bxk</sup>, H-2<sup>sxd</sup>)

References Sastry & Arlinghaus 1991

 Peptides induced T-cell proliferative response to immunizing peptide and to gp160.

**HXB2 Location** gp160 (519–543) **Author Location** Env (519–543)

Epitope FLGFLGAAGSTMGAASLTLTVQARC

Immunogen vaccine

Vector/Type: peptide

Species (MHC) macaque

References Nehete et al. 1993

- Synthetic peptide derived from conserved region of the HIV-1 envelope that stimulates a proliferative response in mice, and in rhesus monkeys.
- Proliferative response to this peptide was observed in 3/3 immunized rhesus monkeys.

**HXB2 Location** gp160 (519–543) **Author Location** Env (519–543)

Epitope FLGFLGAAGSTMGAASLTLTVQARQ

Immunogen HIV-1 infection Species (MHC) human, chimpanzee References Nehete *et al.* 1998b

HIV-infected chimpanzees and HIV-positive patients show positive proliferative responses to multiple peptides from five conserved regions of the HIV-1 Env.

**HXB2 Location** gp160 (547–561)

Author Location gp41 (547–561 IIIB, B10)

Epitope GIVQQQNNLLRAIEA

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (562-576)

Author Location gp41 (562–576 IIIB, B10)

Epitope QQHLLQLTVWGIKQL

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

impaired lymophoproliferative capacity, not viral escape. Gagspecific Th responses were observed more frequently (16%), monly evoke T-cell responses.

**HXB2 Location** gp160 (570–589) **Author Location** gp41 (MN)

Epitope VWGIKQLQARVLAVERYLKD

Epitope name VD20

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DR)

**Assay type** Cytokine production, proliferation, CD4 T-cell Elispot - IFNγ, Intracellular cytokine

staining

References Malhotra et al. 2003

- 92 acute- or early-HIV infected subjects were tested for Envand Gag-specific Th responses. There was an overall low probability of detecting HIV-1-specific Th responses, Env responses were rare and (5%) and only detected very early in infection, probably reflecting a low frequency of response and impaired lymophoproliferative capacity, not viral escape. Gagspecific Th responses were observed more frequently (16%), were found throughout the acute and early infection phases, and predominated after ARV therapy.
- This epitope is one of six Env-specific Th responses detected in one patient 11 days post-infection that were studied in detail. The CD4+ clones showed MHC-restricted cytotoxicity and secreted high levels of cytokines when stimulated. These Th reponses were not detected at subsequent time points, despite the Env specific Th cells being maintained as detected by TCR clonal transcripts, and the epitope sequence being maintained in Env.
- This peptide showed promiscuous binding to DRB1\*0101, DRB1\*1101, DRB1\*1302, DRB1\*0701, DRB1\*0801 DRB4\*0101 DRB5\*01.

**HXB2 Location** gp160 (572–591)

**Author Location** gp41 (572–591)

Epitope GIKQLQARILAVERYLKDQQ

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

References Brown et al. 1995

- This peptide was a good immunogen in BALB/c and CBA mice, producing a strong proliferative response.
- At least one of the four residues GIKQ enhances stimulation, and in CBA mice these residues influence the ability to prime T-cells in vivo.
- QLQARILAVERY stimulated the greatest in vitro T-cell response.
- VERYLKDQQ was the minimal reactive sequence recognized by a T-cell line.

**HXB2 Location** gp160 (576–591)

Author Location gp41 (576–591)

Epitope LQARILAVERYLKDQQ

Immunogen vaccine

Vector/Type: peptide

Species (MHC) mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

References Brown et al. 1995

 This peptide was a poor immunogen in BALB/c and CBA mice used in this experiment, producing a weak proliferative response.

**HXB2 Location** gp160 (578–608) **Author Location** gp41 (585–615 IIIB)

gp160 Helper/CD4+ T-cell epitopes

Epitope ARILAVERYLKDQQLLGIWGCSGKLICTTAV

Immunogen vaccine

Vector/Type: peptide

Species (MHC) mouse

References Goodman-Snitkoff et al. 1990

• Identification of putative Th epitopes that can stimulate an antibody response in peptide immunized mice.

**HXB2 Location** gp160 (579–601) **Author Location** gp41 (579–601)

Epitope RILAVERYLKDQQLLGGIWGCSGK

Immunogen vaccine

Vector/Type: peptide

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>b</sup>)

References Brown et al. 1995

- This peptide was a good immunogen in BALB/c and CBA.
- This peptide produced a strong Th response in both mice strains which was more responsive towards GIKQLQARILAV-ERYLKDQQ and LQARILAVERYLKDQQ than to immunizing peptide.

**HXB2 Location** gp160 (579–604) **Author Location** gp41 (584–609 LAI)

Epitope RILAVERYLKDQQLLGIWGCSGKLIC

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

HXB2 Location gp160 (586–597)

Author Location Env (586–598)

Epitope YLRDQQLLGIWG

Immunogen HIV-1 infection

Species (MHC) human, chimpanzee

References Nehete et al. 1998b

HIV-infected chimpanzees and HIV-positive patients show positive proliferative responses to multiple peptides from five conserved regions of the HIV-1 Env.

HXB2 Location gp160 (586–598) Author Location Env (586–598) Epitope YLRDQQLLGIWGC Immunogen vaccine

Vector/Type: peptide

Species (MHC) macaque, mouse References Nehete *et al.* 1993

- Synthetic peptide derived from conserved region of the HIV-1 envelope that stimulates a proliferative response in mice.
- Proliferative response to this peptide was observed in 1/3 immunized rhesus monkeys, with a weak transient response in the other two.

**HXB2 Location** gp160 (593–604) **Author Location** gp41 (598–609 LAV-1) Epitope LGLWGCSGKLIC Immunogen vaccine

**Species (MHC)** mouse (H2<sup>d</sup>) **References** Schrier *et al.* 1988

 Murine T-dependent B-cell response – 7/29 had a proliferative response to this peptide.

HXB2 Location gp160 (593–604)
Author Location gp41 (593–604 IIIB)
Epitope LGIWGCSGKLIC
Immunogen HIV-1 infection

**Species** (MHC) human **References** Bell *et al.* 1992

• Elicits T-cell proliferation and B cell responses, but only during the asymptomatic phase of HIV infection.

HXB2 Location gp160 (594–603)
Author Location gp41 (594–603 IIIB)
Epitope GIWGCSGKLI
Immunogen HIV-1 infection
Species (MHC) human

References Kelleher et al. 1998b

- Epitope documented as a "previously described' epitope Bell *et al.* [1992], but in Bell *et al.* it was described as gp41(594-603 IIIB), LGIWGCSGKLIC.
- Immunization with a p24-VLP virus-like particle did not significantly impact CD4+ lymphocyte count, viral load, or p24 antibody titre.
- Immunization with p24-VLP did not increase the proliferative response to this gp41 epitope, however, there was a modest, short-lived increased proliferative response to p24.

**HXB2 Location** gp160 (594–604)

Author Location gp41 (consensus)

 ${\bf Epitope} \ {\tt GIWGCSGKLIC}$ 

Immunogen HIV-1 infection

Species (MHC) human

References Mutch et al. 1994

• Core region of peptides that can stimulate proliferative responses from seronegative and seropositive people.

HXB2 Location gp160 (598–609) Author Location gp41 (603–614 LAI) Epitope CSGKLICTTAVP

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

 $\begin{array}{c} \textbf{HXB2 Location} & gp160 \ (604\text{--}615) \\ \textbf{Author Location} & gp41 \ (609\text{--}620 \ LAI) \\ \textbf{Epitope} & \texttt{CTTAVPWNASWS} \end{array}$ 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

**HXB2 Location** gp160 (606–620) **Author Location** gp41 (1035)

**Epitope** TNVPWNASWSNKSLE

Subtype B Immunogen vaccine

Vector/Type: vaccinia prime with gp120 boost Strain: B clade 1035 HIV component: Env Adjuvant: Complete Freund's

Adjuvant (CFA)

Species (MHC) mouse (Class II I Ab)

Assay type T-cell Elispot

**Keywords** epitope processing, vaccine-induced epitopes, escape, TCR usage

References Zhan et al. 2003

 A very narrow Th response was stimulated in C57BL/6 mice vaccinated with vaccinia expressed HIV-1 env clone 1035, to the peptide PKVSFEPIPIHYCAP, located in the C2 region of gp120. The only other peptide recognized using Elispot on Env overlapping peptides to test vaccine responses in the mice was this one: TNVPWNASWSNKSLE, located in gp41.

HXB2 Location gp160 (606–620) Author Location gp41 (UG92005) Epitope TNVPWNASWSNKSLE

Immunogen vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This gp140 epitope of UG92005 (UG, clade D) was recognized by five hybridomas with Vβ usage Vβ 8.1, 14 and not determined one of the Vβ 8.1 was shown to utilize Vα 8.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site

proximity may allow binding to lectins and promote trafficking through processing pathways.

HXB2 Location gp160 (609–616)
Author Location gp41 (consensus)
Epitope PWNASWSN
Immunogen HIV-1 infection

Species (MHC) human

References Mutch et al. 1994

Core region of peptides that can stimulate proliferative responses from seronegative and seropositive people.

HXB2 Location gp160 (611–620) Author Location gp41 (1007, UG92005)

**Epitope** NASWSNKSLE **Immunogen** vaccine

Vector/Type: DNA, protein, vaccinia Strain: B clade 1007, D clade UG92005 HIV component: gp140 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2 IA<sup>b</sup>)

**Keywords** subtype comparisons, epitope processing, TCR usage

References Surman et al. 2001

- This gp41 epitope is conserved in 1007 (US, clade B) and UG92005 (UG, clade D) and was recognized by two hybridomas from two different mice that were vaccinated with different clades the  $V\beta$  usage was  $V\beta$  4 and 14.
- The epitope described here is the region of overlap of two 15 mers that were both able to stimulate IL-2 production from the hybridoma (T[TN]VPWNASWSNKSLE and NASWSNKSLE-QIWNN) the only difference between 1007 and UG92005 for these two proteins is that 1007 has a T and UG92005 has an N in the second position of the first peptide.
- C57BL/6 mice were immunized with a prime-boost strategy involving three HIV-1 Env antigens: Mice were primed with DNA given i.m., 3-4 weeks later boosted with VV, and 3-4 weeks later boosted again with purified protein in Freund's adjuvant.
- The vaccinia construct is a pSC11-based VV vector with the first 38 amino acids contributed by BH10 and the rest of gp120 and gp41 by the vaccine strain, the DNA construct is in the pJW4303 vector with a CMV promotor, and the purified protein is expressed from the pJW4303 vector transfected into CHO-K1 cells.
- Ten days after the final boost, hybridomas were made and tested for IL-2 production using either B6 spleen cells or H-2 IA<sup>b</sup> transfected L cells as targets and Vβ usage was determined.
- Mice were immunized with an Env from either one of two clades: HIV-1 1007, a clade B strain isolated from an individual from Memphis Tennesee, and HIV-1 92UG005, a clade D vaccine isolated from Uganda in 1992 through the WHO.
- 80 unique clonotypes were characterized from six mice.
- H-2 IA<sup>b</sup> restricted T-helper epitopes were concentrated in 5 distinct regions within the Env sequence (2 clonotype responses in V2, 26 in C2, 22 in V3, 23 in V4C4, and 7 in gp41).
- Epitope hotspots tended to be proximal to heavily glycosylated regions of the Env sequence, in exposed, non-helical strands of the protein. The non-uniform localization may be influenced by differential antigen processing and the glycosylation site

proximity may allow binding to lectins and promote trafficking through processing pathways.

**HXB2 Location** gp160 (614–629)

Author Location gp41 (IIIB)

 ${\bf Epitope} \ {\tt WSNKSLEDIWDNMTWC}$ 

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro.
- Peptide priming does not always induce T-cells that recognize whole protein.

**HXB2 Location** gp160 (634–649)

Author Location gp41 (IIIB)

Epitope EIDNYTNTIYTLLEEC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in vitro.
- Peptide priming does not always induce T-cells that recognize whole protein.

**HXB2 Location** gp160 (647–661)

Author Location gp41 (647–661 IIIB, B10)

Epitope EESQNQQEKNEQELL

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

**HXB2 Location** gp160 (650–662)

Author Location gp41 (655–667 LAI)

Epitope QNQQEKNEQELLE

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

**HXB2 Location** gp160 (667–681)

Author Location gp41 (667–681 IIIB, B10)

Epitope ASLWNWFNITNWLWY

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

HXB2 Location gp160 (682-696)

Author Location gp41 (682–696 IIIB, B10)

Epitope IKLFIMIVGGLVGLR

Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

**HXB2 Location** gp160 (724–745)

Author Location gp41 (731–752)

Epitope PRGPDRPEGIEEEGGERDRDRS

Immunogen vaccine

Vector/Type: peptide in cowpea mosaic virus (CPMV) HIV component: gp41 Adjuvant:

Quillaja saponin (Quil-A)

**Species (MHC)** mouse (H-2<sup>k</sup>)

Keywords Th1

References McInerney et al. 1999

- A gp41 peptide was expressed in a cowpea mosaic virus (CPMV) and mice were vaccinated with a purified chimeric particle – out of five adjuvants tested, only Quil A could stimulate anti-gp41 antibodies and an *in vitro* proliferative response.
- The antibodies were predominantly IgG2a, suggesting a Th1 response.

HXB2 Location gp160 (732–744)

**Author Location** gp41 (737–749 LAI)

Epitope GIEEEGGERDRDR

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References Schrier et al. 1989

• Stimulates T-cell proliferation in HIV-infected donors.

HXB2 Location gp160 (780–794)

Author Location gp41 (787-801 IIIB)

**Epitope** RIVELLGRRGWEALK

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse  $(H-2^d, H-2^k, H-2^{t4})$ 

References Hale et al. 1989

• Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

**HXB2 Location** gp160 (780–794)

Author Location gp160 (787–801 IIIB)

Epitope RIVELLGRRGWEALK

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Com-

plete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup>, H-2<sup>s</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- This peptide elicited proliferative responses in cells from B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>), B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), and B10.S(9R) mice (H-2A<sup>s</sup>, E<sup>s</sup>)
- RIVELLGRRGWEALKYWWNLLQYWSQELKNSAVS encompasses several murine Th epitopes including RIVELLGR-RGWEALK and is referred to as a "multideterminant region" or cluster peptide, but the longer peptide only stimulates cells from H-2<sup>k</sup> mice.

**HXB2 Location** gp160 (780–813)

**Author Location** gp160 (787–820 IIIB)

 ${\bf Epitope} \ \ {\tt RIVELLGRRGWEALKYWWNLLQYWSQELKNSA-}$ 

VS

Immunogen HIV-1 infection, vaccine

HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- RIVELLGRRGWEALKYWWNLLOYWSOELKNSAVS encompasses several murine Th epitopes and is referred to as a "multideterminant region" or cluster peptide.
- Six multideterminant region cluster peptides were evaluated Th responses in different MHC/HLA backgrounds after vaccination of mice with gp160, or in infected people.
- This cluster peptide elicited proliferative responses in cells from only B10.BR mice (H-2Ak, Ek), and not from B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), B10.A(5R) mice (H-2A<sup>b</sup>, E<sup>b</sup>), or B10.S(9R) mice (H-2A<sup>s</sup>, E<sup>s</sup>)
- IL-2 production in response to this peptide was observed in 59% (17/29) of asymptomatic HIV-infected individuals.

HXB2 Location gp160 (794–808) Author Location gp41 (801–815 IIIB) Epitope KYWWNLLQYWSQELK

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse  $(H-2^k)$ References Hale et al. 1989

• Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

**HXB2 Location** gp160 (794–808) Author Location gp160 (801–815 IIIB) Epitope KYWWNLLQYWSQELK

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup>, H-2<sup>s</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

- This peptide elicited proliferative responses in cells from B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>), B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), and B10.S(9R) mice (H-2As, Es)
- RIVELLGRRGWEALKYWWNLLQYWSQELKNSAVS encompasses several murine Th epitopes including KYWWNL-LQYWSQELK and is referred to as a "multideterminant region" or cluster peptide, but the longer peptide only stimulates cells from H-2<sup>k</sup> mice.

**HXB2 Location** gp160 (799–813) Author Location gp160 (806–820 IIIB) **Epitope** LLQYWSQELKNSAVS

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup>, H-2<sup>s</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

• This peptide elicited proliferative responses in cells from B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>), B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), and B10.S(9R) mice (H-2A<sup>s</sup>, E<sup>s</sup>)

Vector/Type: protein Strain: B clade IIIB • RIVELLGRRGWEALKYWWNLLOYWSOELKNSAVS encompasses several murine Th epitopes including LLOY-WSQELKNSAVS and is referred to as a "multideterminant region" or cluster peptide, but the longer peptide only stimulates cells from H-2k mice.

> HXB2 Location gp160 (799–813) Author Location gp41 (806–820 IIIB) **Epitope** LLQYWSQELKNSAVS

> > Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse  $(H-2^k, H-2^d, H-2^{t4})$ 

References Hale et al. 1989

· Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

**HXB2 Location** gp160 (799–813) Author Location gp41 (806–820 IIIB) **Epitope** LLQYWSQELKNSAVS

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>d</sup>, H-2<sup>t4</sup>)

References Hale et al. 1989

• Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

HXB2 Location gp160 (814-829)

**Author Location** gp41 (IIIB)

**Epitope** WLNATAIAVTEGTDRC

Immunogen in vitro stimulation or selection

Species (MHC) human

References Manca et al. 1995b

- Peptide stimulation of PBMC from non-infected individuals in
- Peptide priming does not always induce T-cells that recognize whole protein.

HXB2 Location gp160 (821–835)

Author Location gp41 (828-842 IIIB)

Epitope AVAEGTDRVIEVVQG

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>k</sup>)

References Hale et al. 1989

• Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

HXB2 Location gp160 (821–835) Author Location gp160 (828–842 IIIB) Epitope AVAEGTDRVIEVVQG

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>b</sup>, H-2<sup>s</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

• This peptide elicited proliferative responses in cells from B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>), B10.A(5R) mice (H-2A<sup>b</sup>, E<sup>b</sup>), and B10.S(9R) mice  $(H-2A^s, E^s)$ 

AVAEGTDRVIEVVQGAYRAIRHIPRRIRQGLER encompasses several murine Th epitopes including AVAEGT-DRVIEVVQG and is referred to as a "multideterminant region" or cluster peptide.

**HXB2 Location** gp160 (821–838) **Author Location** gp41 (827–843)

Epitope YVAEGTDRVIEVVQGACR

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression **References** Caruso *et al.* 1997

- As HIV-1-infected individuals progress to disease, T-cells show reduced ability to proliferate in response to HIV antigen, but retain the ability to express the activation antigens CD25 and CD71.
- The ability to express activation markers in response to HIV is retained, but the response to tetanus toxoid recall antigen is lost.
- This study investigated CD25 and CD71 expression in PBMC from patients at various stages of progression, measuring the response to *in vitro* stimulation by peptide cocktail containing four antigenic Env peptides, or p17 and p24.

**HXB2 Location** gp160 (821–853)

Author Location gp160 (828–860 IIIB)

Epitope AVAEGTDRVIEVVQGAYRAIRHIPRRIRQGLER

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) human, mouse (H-2<sup>k</sup>, H-2<sup>b</sup>, H-2<sup>s</sup>, H-2<sup>d</sup>)

**References** Berzofsky et al. 1991b; Berzofsky et al. 1991a

- AVAEGTDRVIEVVQGAYRAIRHIPRRIRQGLER encompasses several murine Th epitopes and is referred to as a "multideterminant region" or cluster peptide.
- Six multideterminant region cluster peptides were evaluated for Th responses in different MHC/HLA backgrounds after vaccination of mice with gp160, or in infected people.
- This cluster peptide elicited proliferative responses in cells from all four MHC types tested: B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>), B10.D2 mice (H-2A<sup>d</sup>, E<sup>d</sup>), B10.A(5R) mice (H-2A<sup>b</sup>, E<sup>b</sup>), and B10.S(9R) mice (H-2A<sup>s</sup>, E<sup>s</sup>)
- IL-2 production in response to this peptide was observed in only 8% (1/12) of asymptomatic HIV-infected individuals.

**HXB2 Location** gp160 (827–835)

Author Location gp41 (834–842 IIIB)

Epitope DRVIEVVQG Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species** (MHC) mouse (H-2<sup>k</sup>) **References** Hale *et al.* 1989

Suggested H-2<sup>k</sup> epitope based on region of overlap.

HXB2 Location gp160 (827–841) Author Location gp160 (834–848 IIIB) Epitope DRVIEVVQGAYRAIR

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>b</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

 This peptide elicited proliferative responses in cells from B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>) and B10.A(5R) mice (H-2A<sup>b</sup>, E<sup>b</sup>)

**HXB2 Location** gp160 (827–841) **Author Location** gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4
Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>i5</sup>) **References** Hale *et al.* 1989

- Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.
- Called Th4.1 and TH4.

**HXB2 Location** gp160 (827–841) **Author Location** gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4
Immunogen vaccine

*Vector/Type:* peptide prime with protein boost *Strain:* B clade IIIB *HIV component:* gp160

Species (MHC) macaque

References Hosmalin et al. 1991

- Peptide priming to induce T-cell help enhances antibody response to gp160 immunization.
- Called Th4.1 and TH4.

**HXB2 Location** gp160 (827–841)

**Author Location** gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1997

used in a study of the influence of pentoxifylline on HIV specific T-cells.

**HXB2 Location** gp160 (827–841)

Author Location gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4

Immunogen

Species (MHC) human

References Pinto et al. 1995

- CTL activity analyzed in parallel with Th reactivity in exposed but uninfected health care workers.
- Called Th4.1 and TH4.

**HXB2 Location** gp160 (827–841)

Author Location gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1991a

- Peptides stimulate Th cell function and CTL activity in similar patient populations.
- Called Th4.1 and TH4.

**HXB2 Location** gp160 (827–841)

Author Location gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4
Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp160

Species (MHC) human

References Clerici et al. 1991b

- Immunizing uninfected individuals with rgp160 results in stronger Th response than does natural infection.
- Called Th4.1 and TH4.

**HXB2 Location** gp160 (827–841)

Author Location gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4

Immunogen

Species (MHC) human

References Clerici et al. 1992

- Cell-mediated immune response to HIV-1 peptides in HIV-1 exposed seronegative men.
- Called Th4.1 and TH4.

**HXB2 Location** gp160 (827–841)

Author Location gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4

Immunogen HIV-1 infection

Species (MHC) human

References Clerici et al. 1989

- IL-2 production detection of Th lymphocytes from asymptomatic HIV-positive individuals.
- · Called Th4.1 and TH4.

**HXB2 Location** gp160 (827–841)

Author Location gp41 (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4

Immunogen HIV-1 infection

Species (MHC) human

References Kaul et al. 1999

- Kenyan sex workers that remained seronegative were found to frequently have HIV-env peptide specific Th responses detected by an IL-2 assay (11/20 cases) and mucosal genital tract anti-HIV IgA (16/21 cases)
- Helper epitopes used in this study were noted to be previously described Clerici et al. [1989], and were not explicitly described in Kaul et al. [1999]

HXB2 Location gp160 (827-841)

Author Location gp41

Epitope DRVIEVVQGAYRAIR

Epitope name TH4, Th4.1

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC) human

**Keywords** subtype comparisons, responses in children, mother-to-infant transmission

References Kuhn et al. 2001a

- In a S. African perinatal transmission study, 33% (33/86) of cord blood samples from infants with seropositive mothers produced T-helper responses (measured by a bioassay measuring IL2 production in a murine cell line and confirmed with a proliferation assay) against a peptide cocktail containing Th epitopes P18 MN, P18 IIIB, T1, T2, and TH4.
- The mothers were predominantly infected with subtype C, but the T help response was detectable in a number of cord blood samples despite using peptides based on B subtype reagents.
- 3/33 infants with cord blood T help responses to Env were infected in utero, 2/33 were lost to follow up, and 28/33 were not infected. 6/53 of the infants with cord blood that was unresponsive to Env peptide stimulation were infected before delivery, and 8/47 contracted HIV intrapartum or via breastfeeding.
- Measurable HIV specific T help responses elicited in the immunologically immature newborn, possibly in response to in utero exposure, are associated with a protective natural immunity that helps block mother-infant transmission of HIV-1.

**HXB2 Location** gp160 (827–841)

**Author Location** Env (834–848 IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4.1

Immunogen HIV-1 infection, HIV-1 exposed seronegative

Species (MHC)

Assay type Cytokine production

Keywords mother-to-infant transmission

References Clerici et al. 1993a

- Cord blood samples in 8/23 infants with HIV+ mothers showed IL-2 production in response to peptides from HIV-1 gp1260, demonstrating that Th cell priming to HIV env determinants can occur in utero. HIV-specific Th immunity in utero may be protective, as none of the 8 with HIV-1 specific Th activity became infected, while 3/15 infants with no detectable Th activited were infected.
- PBL from 10/21 of the mother showed HIV-1 specific Th activity through IL-2 production in response to Env peptides.

**HXB2 Location** gp160 (827–841)

**Author Location** Env (IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4.1

Subtype B

Immunogen HIV-1 exposed seronegative

Species (MHC)

Assay type Cytokine production

References Clerici et al. 1994a

 Six of eight HIV-exposed health care workers had transient HIV-specific T-helper responses after percutaneous exposure to HIV, responding to two or more Env peptides among the set T1, T2, Th4.1, P18IIIB and P18MN. Responses were lost after 12-56 weeks. The HIV-specific Th responses occurred without seroconversion or PCR evidence for infection. • Six of the eight HIV-exposed individuals responded to two • Viral isolates (gp160) from 16 vertically HIV-1 infected chilor more peptides, whereas, only one individual exposed to HIV-negative blood responded to one peptide.

HXB2 Location gp160 (827–841) **Author Location** HIV-1 (IIIB)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4.1 Subtype B

Immunogen HIV-1 infection

Species (MHC)

Assay type Cytokine production References Clerici et al. 1994b

• IL-10-specific mRNA was upregulated in PBMC from asymptomatic, HIV-infected (HIV+) patients, and was particularly high in those with severely compromised Th cells function. Th response to HIV peptides in vitro could be restored by IL-10

HXB2 Location gp160 (827-841) **Author Location** Env (834–848)

Epitope DRVIEVVQGAYRAIR

Epitope name TH4-1

Immunogen HIV-1 infection

Species (MHC) human

Assay type proliferation

Keywords responses in children, mother-to-infant trans-

mission

References Kuhn et al. 2001b

- Th proliferative responses in cord blood at delivery to a cocktail of HIV envelope peptides were infrequent (1/41) among infants whose HIV-positive mothers received antiretroviral prophylaxis at delivery to prevent transmission. Responses were 10 times more frequent among infants of HIV-seropositive women who had no antiretroviral treatment (7/29). Reductions of HIVspecific responses occurred despite persistance of detectable HIV RNA in the mothers at delivery.
- The reduction of Th responses in newborns raises the possibility that anti-retrovial exposure during pregnancy may block subsequent immune protection. The authors point two relevant citations that showed ARV exposed infants had a more rapid course of progression (Kuhn et al., JID 182:104 (2000)), but were at no greater risk of infection due to subsequent breast feeding (Ditrame et al., Lancet 354:2050 (1999)).

HXB2 Location gp160 (827–841)

Author Location Env (gp160) (421–436)

Epitope DRVIEVVGQAYRAIR

Epitope name TH4.1

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Country South Africa. Assay type proliferation

Keywords responses in children, variant cross-

recognition or cross-neutralization

References Meddows-Taylor et al. 2004

- dren (8 T-helper cell Env responders and 8 non-responders) were analyzed for variation after comparison with Env peptide sequences (conserved regions T1, T2 and TH4.1, hypervariable regions P18 MN and P181 IIIB) used for in vitro stimulation.
- No correlation between the age, clinical category (mild or severe), HIV-1 viral load and the degree of epitope variation was established.

HXB2 Location gp160 (827-853)

**Author Location** Env (HIV-1 IIIB)

Epitope DRVIEVVQGAYRAIRHIPRRIRQGLER

Subtype B

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB, SIV HIV component: Env, Gag, Pol Adjuvant: E. coli mutant heat labile enterotoxin

(LT-R72), Montanide (ISA 51)

Species (MHC) macaque

Assay type proliferation Keywords mucosal immunity References Belyakov et al. 2001

- · Different HIV strains were used for different regions: env HIV-1 IIIB, gag SIV, pol SIV
- Intrarectal vaccination with a Th and CTL peptide vaccine provided better protection against intrarectal challenge with pathogenic SHIV-Ku1 than subcutaneous administered vaccine. In some animals after the initial viremia, viral loads were diminished to undetectable levels in the blood and intestine, and CD4+ T cells were better preserved.
- The CD4 T-cell proliferative response correlated with the level of the CTL response.

HXB2 Location gp160 (829–843)

Author Location gp160 (836–850 IIIB)

Epitope VIEVVQGAYRAIRHI

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>b</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

• This peptide elicited proliferative responses in cells from B10.BR mice (H-2A<sup>k</sup>, E<sup>k</sup>) and B10.A(5R) mice (H-2A<sup>b</sup>, E<sup>b</sup>)

HXB2 Location gp160 (834–841)

Author Location gp41 (841–848 IIIB)

Epitope QGAYRAIR Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

Species (MHC) mouse (H-2<sup>i5</sup>)

References Hale et al. 1989

• Suggested H-2<sup>k</sup> epitope based on region of overlap.

HXB2 Location gp160 (834–848)

Author Location gp41 (841-855 IIIB)

Epitope QGAYRAIRHIPRRIR

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>t4</sup>, H-2<sup>i5</sup>)

References Hale et al. 1989

 Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

HXB2 Location gp160 (834–848) Author Location gp160 (841–855 IIIB) Epitope QGAYRAIRHIPRRIR

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** mouse (H-2<sup>k</sup>, H-2<sup>b</sup>, H-2<sup>d</sup>, H-2<sup>s</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

• This peptide elicited proliferative responses in cells from B10.BR mice (H-2A $^k$ , E $^k$ ), B10.A(5R) mice (H-2A $^b$ , E $^b$ ), B10.D2(H-2A $^d$ , E $^d$ ), and B10.S(9R) mice (H-2A $^s$ , E $^s$ )

HXB2 Location gp160 (839–848) Author Location gp41 (846–855 IIIB) Epitope AIRHIPRRIR Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>t4</sup>) **References** Hale *et al.* 1989

• Suggested H-2<sup>d,t4</sup> epitope based on region of overlap.

HXB2 Location gp160 (839–853) Author Location gp41 (846–860 IIIB) Epitope AIRHIPRRIRQGLER

Immunogen vaccine

Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>, H-2<sup>t4</sup>) **References** Hale *et al.* 1989

 Six multideterminant helper T-cell regions are recognized by mice of three or four MHC types.

HXB2 Location gp160 (839–853) Author Location gp160 (828–842 IIIB) Epitope AIRHIPRRIRQGLER

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp160 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (MHC)** human, mouse (H-2<sup>k</sup>, H-2<sup>b</sup>, H-2<sup>s</sup>)

References Berzofsky et al. 1991b; Berzofsky et al. 1991a

• This peptide elicited proliferative responses in cells from B10.BR mice (H-2A $^k$ , E $^k$ ), B10.A(5R) mice (H-2A $^b$ , E $^b$ ), and B10.S(9R) mice (H-2A $^s$ , E $^s$ )

HXB2 Location gp160 (842-856)

Author Location gp41 (842–856 IIIB, B10)

Epitope HIPRRIRQGLERILL Immunogen HIV-1 infection

Species (MHC) human

References Wahren et al. 1989b; Wahren et al. 1989a

• 12 gag and 18 env peptides were identified that could commonly evoke T-cell responses.

## III-B-15 Env Helper/CD4 + T-cell epitopes

**HXB2 Location** Env **Author Location** gp120

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env, Gag, Pol Adjuvant: IFNγ, IL-2, IL-4

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1

References Kim et al. 2000

• Co-stimulatory molecules co-expressed with an HIV-1 immunogen in a DNA vaccine used to enhance the immune response – co-expression of Th1 cytokine IFN $\gamma$  drove Th1 immune responses and enhanced CTL responses.

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: gp160

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords Th1, Th2

References Shirai et al. 2001

Helicobacter pylori induces Th1 responses early, but predominantly Th2 responses later in infection (at 6 weeks) – differentiation of HIV-1 gp160 CD4+ help and CD8+ CTL effector cells in response to HIV gp160-vaccinia vaccination is impaired in BALB/c mice infected with H. pylori.

**HXB2 Location** Env

Author Location gp120 (V3) and p24 (IIIB, MN, BH10)

Epitope Subtype A, B Immunogen vaccine

Vector/Type: virus-like particle (VLP) Strain: A clade UG5.94UG018, B clade IIIB

HIV component: Gag, gp120

**Species (MHC)** mouse (H-2<sup>d</sup>)

**Keywords** subtype comparisons **References** Buonaguro *et al.* 2002

- Different HIV strains were used for different regions: gp120 A clade UG5.94UG018; Gag HIV-1 IIIB
- BALB/c mice were given intraperitoneal immunization in the absence of adjuvants with virus-like particles (VLPs) expressing recombinant subtype A gp120 and Pr55gag.
- High dose-independent humoral responses were elicited against both gp120 and p24 peptides, and CTL responses were observed against target cells carrying vaccinia expressed gp120 and Gag.
- Recombinant rgp120 (clade B, MN) induced T cell proliferative responses in vitro from vaccinated animals.

**HXB2 Location** Env

Author Location gp160 (IIIB)

**Epitope** 

Immunogen vaccine

Vector/Type: peptide, protein Strain: B clade IIIB HIV component: gp160, V3 Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

**Species (MHC)** mouse (H2<sup>d</sup>)

Keywords Th1, Th2

References Morris et al. 2000

- Mice were intranasally immunized with 20 ug of HIV-gp160 and 5 ug of peptide E7 (RIHIGPGRAFYAARK) with the adjuvant LT(R192G), a heat-labile enterotoxin produced by E.
- Adjuvant LT(R192G) was required for stimulation of antigenspecific IgG1, IgG2 antibodies, and Th1 and Th2 cytokines responses to gp160, and peptide-specific CTL responses.
- Increased IFNγ, IL-10 and IL-6 cytokine production specific to gp160 was measured with co-immunization of gp160 with LT(R192G)

**HXB2 Location** Env

Author Location gp160 (IIIB)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade IIIB HIV component: gp160, Rev Adjuvant: Br-cAMP

**Species (MHC)** mouse (H2<sup>d</sup>)

Keywords Th1

References Arai et al. 2000

- The CMV promotor responds to the intracellular level of cAMP, and 8 Br-cAMP can increase transgene expression so it was coadministered with a CMV-based DNA vaccine both intranasally and intramuscularly.
- 8 Br-cAMP increased serum IgG responses, HIV-specific CTL, DTH and Th1 responses, and IgA in the intranasal vaccination.
- · A CAT assay study showed adjuvant effect was due to CMV promotor activation.

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB HIV component: gp120, gp160

Species (MHC) mouse

Keywords Th1

References Shiver et al. 1997

- DNA vaccinations of BALBc mice with a gp120 or gp160 DNA vaccine elicited a strong T-cell proliferative response with Th1-like secretion of  $\gamma$  interferon and IL-2, with little or no IL-4, as well as antigen specific gp120 Abs.
- · An intramuscular route of inoculation gave a stronger proliferative response than intradermal.
- A proliferative response could be detected in all lymph tissues tested: spleen, PBMC, and mesenteric, iliac, and inguinal lymph nodes.

**HXB2 Location** Env Author Location gp120

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Gag, gp160, Pol Adjuvant: CD86

Species (MHC) mouse

References Kim et al. 1997d

• A gp160 DNA vaccine, when delivered in conjunction with the plasmid encoding the co-stimulatory molecule CD86, gives an increase in the proliferative responses to gp120 in mice.

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Immunogen

Species (MHC) human

References De Berardinis et al. 1997

· Sequences flanking helper T-cell immunogenic domains can be important for immunogenicity.

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Rosenberg et al. 1997

• A strong proliferative response to p24 and gp160 was found in a healthy long term survivor.

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) macaque

Keywords Th1, Th2

References Kent et al. 1997b

- · Macaca nemestrina can be infected with HIV, and clear the infection within 6 months, so it is of interest to examine their initial immune response.
- A strong proliferative response against gp160 with IL-4 production, indicating a Th2 response, was found with 4 weeks of infection.
- The gp160 proliferative response by 8 weeks produces both IL-4 and  $\gamma$  interferon, indicating both Th1 and Th2 responses.

**HXB2 Location** Env

Author Location gp120 (HXBc2)

Epitope

Immunogen vaccine

Vector/Type: DNA prime with gp160 boost Strain: B clade HXBc2 HIV component:

gp160

Species (MHC) macaque

References Letvin et al. 1997

- · Vaccination of Macaca mulatta (rhesus monkeys) with a HXBc2 env DNA prime and a protein boost elicited a T-cell proliferative response, a CTL response, and type-specific neutralizing antibodies.
- · Vaccinated animals challenged with SHIV-HXB2 were protected from infection.

**HXB2 Location** Env

Author Location gp120 (MN)

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA Strain: B clade MN

HIV component: Env, Rev

Species (MHC) human

References MacGregor et al. 1998

- An HIV DNA env and rev vaccine given to 15 asymptomatic HIV+ individuals at three different dosages, 30, 100 or 300 μg, was safe.
- All three groups showed an increased proliferative response after vaccination.

**HXB2 Location** Env

**Author Location** Env

Epitope

Immunogen

Species (MHC) human

References Mazzoli et al. 1997

- Study of HIV-specific immunity in seronegative partners of HIV-positive individuals Env peptides could stimulate IL-2 production in 9/16 HIV-exposed seronegative individuals, and only 1/50 low-risk controls.
- Exposed-uninfected produced more IL-2 and less IL-10 then HIV-infected individuals.
- 8/9 of those whose PBMC produce IL-2 in response to Env peptides had concomitantly detected urinary or vaginal tract anti-HIV IgA.

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Plana et al. 1998

 Patients from later stages of infection given HAART do not show restoration of HIV-1 specific Th proliferative responses.

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Kelleher et al. 1998a

• Env and gag Th epitopes were pooled and used to test Th proliferative responses after IL2 therapy – while IL2 therapy causes an increase in CD4+ lymphocyte count, it does not increase HIV-1 specific proliferative responses.

HXB2 Location Env

Author Location gp160

**Epitope** 

Immunogen HIV-1 infection, vaccine

*Vector/Type:* protein *HIV component:* 

gp160

Species (MHC) human

References Ratto-Kim et al. 1999

 Vaccinations with rgp160 did not enhance Th immunoproliferative responses in individuals who were immunized every 2 months for 5 years starting early in infection.

HXB2 Location Env

**Author Location** gp160

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: protein HIV component:

gp160

Species (MHC) human

Keywords subtype comparisons

References Leandersson et al. 2000

- 27 HIV subtype B, 4 subtype C, 2 D and one of each subtype E, F, G infected individuals were either given rgp160 B clade immunizations or placebo. All rgp160 immunized individuals showed increased proliferation responses to the B clade immunizing antigen rgp160.
- gp120 was prepared from A, B, C, D, and E subtype virions and used as antigenic stimulus 7 of 10 tested individuals responded to native gp120 from at least one additional subtype in addition to B subtype, while a placebo recipient did not respond to any gp120.
- This study shows that cross-subtype HIV-specific T-cell proliferative responses can be stimulated in patients already infected with another HIV-1 subtype all immunized subjects could respond to the subtype B immunogen, but many developed responses to at least one more subtype.

**HXB2 Location** Env

**Author Location** gp160 (MN)

**Epitope** 

Immunogen vaccine

Vector/Type: gp160 prime with gp120 boost Strain: B clade MN HIV component: gp120,

gp160

Species (MHC) human

Keywords Th1, Th2

References Gorse et al. 1999a

 Helper T-cell memory responses were induced by MN rgp160 as measured by proliferation and Th1 and Th2 cytokine release

 this response could be boosted by MN rgp120.

HXB2 Location Env

Author Location gp120

**Epitope** 

Immunogen vaccine

Vector/Type: fowlpoxvirus, ISCOM Strain: B clade SF2 HIV component: gp120

Species (MHC) macaque

Keywords Th1, Th2

References Heeney et al. 1998b

- Vaccinated monkeys with the highest level of Th1 and Th2
  responses and the highest levels of NAbs were protected against
  a SHIV SF13 challenge the ISCOM strategy gave more potent
  anti -gp120 responses than the Fowl pox strategy.
- When animals were challenged 4 months after boost, those that maintained high levels of HIV-1 specific IFNγ responses, indicative of a Th 1 response, were still protected.

HXB2 Location Env Author Location (IIIB) Epitope

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA Strain: B clade IIIB

HIV component: Env, Rev

Species (MHC) human

References Boyer et al. 1999

- A DNA vaccine containing env and rev was tested for safety and immune response in 15 HIV+ asymptomatic individuals.
- Enhanced proliferative activity and higher levels of MIP-1 alpha were detected in multiple study subjects.

HXB2 Location Env Author Location Env Epitope Immunogen vaccine

> Vector/Type: vaccinia Strain: B clade IIIB HIV component: gp160 Adjuvant: GM-

CSF/ENV chimera

Species (MHC) mouse

References Rodríguez et al. 1999

 A chimeric GM-CSF-env antigen expressed in a vaccinia vector elicits a higher HIV-specific env cellular immune response than when native env is used.

HXB2 Location Env Author Location Env (LAI)

Epitope Subtype B Immunogen vaccine

Vector/Type: DNA prime with vaccinia boost Strain: B clade LAI HIV component: Env,

Gag

Species (MHC) macaque Keywords Th1, Th2 References Kent *et al.* 1998

- Priming with an HIV-DNA vaccine and boosting with a vaccinia construct induced greater levels of HIV T-cell immunity than either vaccine alone.
- The proliferative response to Env and Gag after the DNA vaccination had a mean SI of 1.5-4, but after boosting with rHIV-fowlpox virus, there was a 6-17 fold increase in the mean SI for HIV Gag and Env. The T help response happened despite a fall in antibody titers, suggesting that the Th response was primarily Th1, not Th2. The CTL response was also enhanced.

HXB2 Location Env Author Location gp120 Epitope

Immunogen vaccine

Vector/Type: DNA, protein, virus-like particle

(VLP), ISCOM

Species (MHC) macaque Keywords Th1, Th2

References Heeney et al. 1999

- Ten different vaccine strategies were evaluated for their ability
  to protect from infection in a rhesus macaque model using a
  non-pathogenic SHIV challenge. Protection correlated with
  the magnitude of NAb responses, beta-chemokines, and a balanced Th response. DNA, protein+adjuvant, VLP and ISCOM
  vaccines were tested.
- HIV-1/ISCOMS gave the highest NAb titers, Th1 and Th2 responses, was the only vaccine formulation tested with a detectable CTL response, and gave enhanced beta-chemokine production.

**HXB2 Location** Env

Author Location gp160 (MN)

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp160

Species (MHC) human

References Kundu et al. 1998a

- This study followed 10 HLA-A2 asymptomatic HIV+ individuals as they received MN gp160 vaccinations over a two year period.
- There was an increased lymphoproliferative response but this did not impact viral load or CTL response.

**HXB2 Location** Env

Author Location gp120 (SF2)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA, protein, ISCOM Strain: B clade SF2 HIV component: gp120 Ad-

juvant: MF59

Species (MHC) macaque

References Verschoor et al. 1999

- 16 rhesus Macaques were vaccinated with either an epidermal SF2 gp120 DNA vaccine, rgp120 with a MF59 adjuvant, or rgp120 incorporated into ISCOMs.
- DNA vaccination elicited a weak Th type 1 response and low antibody response, rgp120/MF59 triggered a strong antibody response, and rgp120/ISCOM induced both kinds of Th cells, and a strong humoral response.
- Animals were challenged with SF13 SHIV. Early induction of Th type 1 and type 2 responses with the rgp120/ISCOM vaccine provided the most effective immunity, protecting from infection.

**HXB2 Location** Env

**Author Location** Env (MN)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Strain: B clade MN HIV component: Env, Gag, Pol Adjuvant: CD80, CD86

Species (MHC) mouse

References Kim et al. 1998

Co-stimulatory molecules co-expressed with an HIV-1 immunogen in a DNA vaccine used to enhance the immune response – co-expression of CD86, but not CD80, dramatically increased both HIV Env and Gag/Pol specific CTL and Th proliferative responses.

**HXB2 Location** Env

Author Location Env (LAI, MN)

**Epitope** 

Immunogen vaccine

Vector/Type: canarypox Strain: B clade LAI, B clade MN HIV component: Gag, gp120, gp41, Protease

Species (MHC) human

References Salmon-Ceron et al. 1999

 A live attenuated canarypox vector expressing MN gp120 and LAI gp41/gag/protease could induce CTL and a lymphoproliferative response in healthy uninfected volunteers.

HXB2 Location Env Author Location Env Epitope Immunogen vaccine

Vector/Type: DNA Strain: ZF1 HIV com-

ponent: complete genome

Species (MHC) macaque

References Akahata et al. 2000

- Rhesus macaques were vaccinated by i.m. injection with naked plasmid DNA carrying an HIV-1 complete genome vaccine, strain ZF1, with a mutated zinc finger in the nucleocapsid to prevent packaging.
- Env and Gag specific CTL but no antibody responses were induced in 2/4 vaccinated monkeys (MM145 and MM153)
- 2/4 monkeys (MM146 and MM143) produced antibodies against p24 and/or gp160, but no CTL response was detected.
- PBMC from all vaccinated monkeys produced IFN $\gamma$ , in response to HIV-1 gp160, indicating a Th response this response was 5 times higher in MM145, the animal with the strongest CTL response.
- 4 weeks post-challenge with SHIV NM-3rN plasma viral loads of both MM145 and MM153 (with a homologous Env) decreased to near or below the detection limit.
- 6-8 weeks post-challenge with SHIV NM-3rN plasma viral loads of both MM146 and MM143 decreased near or below the detection limit.

**HXB2 Location** Env

Author Location gp120 (W6.ID)

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

References Zhang et al. 2001b

• T-helper cell proliferative responses to HIV p24, p55 and gp120 were tested in 27 patients with HIV infection – vigorous responses directed at Gag were detected in ten patients, but an Env specific response was detected in only one patient.

HXB2 Location Env Author Location gp160 Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART **References** Blazevic *et al.* 2000  Prolonged viral suppression resulting from potent antiretroviral therapy did not allow an HIV T helper response to p24 or gp160, but Th proliferative responses to influenza, alloantigen, and PHA did develop in many HIV+ patients, and asymptomatic patients had stronger and more frequent Th response recovery than AIDS patients.

**HXB2 Location** Env **Author Location** gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART

References Oxenius et al. 2000

 Patients who started therapy at acute HIV infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.

**HXB2 Location** Env **Author Location** gp120

**Epitope** 

Immunogen vaccine

Vector/Type: canarypox prime with gp120

boost HIV component: gp120

Species (MHC) human

Keywords Th1, Th2

References Sabbaj et al. 2000

- Proliferative responses in PBMC of uninfected individuals that were vaccinated with canarypox vector expressing HIV-1 antigens (ALVAC-HIV) and boosted with a recombinant gp120 subunit vaccine gave a Th1 and Th2 proliferative response upon stimulation with HIV-1 Env.
- All vaccinees produced IFN

   and IL10, most also produced IL-2, IL-6, IL-4 and IL-5.

HXB2 Location Env

**Author Location** gp120

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp120

Species (MHC) human

Keywords Th1

References Hladik et al. 2001

- 16/29 HIV-1 infected and 24/30 vaccinated individuals had DTH reactions within 48 hours after an intradermal rec gp120 injection. Of nine DTH positive individuals, none had detectable proliferative responses. Thus skin testing may be a sensitive way to identify people with Th recall responses to vaccines, or in the absence of lymphoproliferaion.
- No 48 hour DTH responses were detected among uninfected volunteers, although 10/35 (40%) of the high risk and 11/32 (34%) of the low risk individuals developed an induration resembling DTH after 7-12 days, that may be indicative of primary induction of HIV-1 specific Th1-immunity.

HXB2 Location Env Author Location gp120 Epitope

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression, Th1, Th2

References Wilson et al. 2000b

- Dysfunction of HIV-1 specific proliferative responses, but not responses to other antigens, is evident in HIV-1 progressive disease.
- Vigorous HIV-1 specific responses to p24, Nef and gp120 with SI between 8-99 were seen in 6/7 long term non-progressors (LTNP), the seventh had a borderline responses. IL-2 production was seen in all cases, and IL-4 production was also evident many responses.
- None of the progressors (0/5) had HIV-1 specific proliferative responses, or IL-2 or IL-4 induction.
- Non-HIV antigens (cytomegalovirus, PPD, Staphylococcus enterotoxin B, tetanus toxoid) gave similar responses in HIV-1 + LTNP, progressors, and HIV-1 controls.

HXB2 Location Env Author Location gp160 Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** rate of progression, Th1 **References** Kalams *et al.* 1999a

- The strength of p24 specific Gag proliferative responses (SIs) were inversely correlated with viral load in 21 ARV naive patients. The responses were Th1, IFNgamma producing.
- Proliferative responses against gp160 were rarely observed (only 4 cases).

HXB2 Location Env Author Location Env Epitope Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade MN HIV component: Env, Rev Adjuvant: Bupivacaine

Species (MHC) human

**Keywords** early-expressed proteins **References** MacGregor *et al.* 2002

- A phase I clinical trial of a HIV-1 Env and Rev DNA vaccine
  with a CMV promoter was conducted and Th proliferative,
  CTL and Elispot responses monitored. The construct was
  modified for safety and included no LTRs or packaging signals.
  The vaccine strategy was safe, and elicited strong CD4-T cell
  responses, but not CD8 T-cell responses. Rev elicited strong
  Th responses, and is a early produced protein so may confer
  advantages.
- With a 300 ug dose, 4/6 individuals had a lymphocyte proliferation (LP) responses to gp120, 3/6 to Rev.
- With a 1000 ug dose, 4/6 individuals had a LP and 2/6 had IFNgamma Elispot responses to gp160; 3/6 had LP, and 4/6 had IFNgamma Elispot responses to Rev.

 No responses to three specific CTL epitopes were observed by Elispot in individuals with appropriate HLA. Some cytotoxic activity against whole protein was observed that was CD4+ T-cell mediated.

HXB2 Location Env
Author Location
Epitope
Immunogen HIV-1 infection
Species (MHC) human
Keywords HAART, ART

References Clerici et al. 2002b

- Specific immunity was compared in a two-year study of chronically HIV-1 infected i) HAART-naive patients who were not progressing and had strong immune responses, ii) newly treated patients followed for 24 months after initiation of HAART, iii) and long-term HAART patients who had been on HAART at least 12 months prior to the study.
- HAART naïve patients had strongest proliferative responses at time zero, but long-term HAART patients the most significant increase in specific responses over the two year study period against HIV-1 gp160, influenza, and Candida. Similarly, IL-2 and IFN $\gamma$  production in responses to gp160 was highest in the naïve group at time zero, but increased the most in the long-term HAART treated patients.
- Short-term HAART patients showed a significant improvement in their CD4+ T cell count and a reduction of plasma viremia, and had augmented IL-7 production, which was slightly reduced in long-term HAART patients.

Author Location gp160
Epitope
Immunogen HIV-1 infection
Species (MHC) human
Keywords HAART, ART
References Palmer et al. 2002

**HXB2 Location** Env

- CD4+ T cell proliferative responses from 33 HIV-1 infected patients with HAART suppression were compared to 19 patients with active viral replication (HAART failures and HAART naive). Patients with HAART suppression showed stronger p24- and p66-specific proliferative responses compared to patient groups with active HIV-1 replication, suggesting active viral replication *in vivo* specifically reduces proliferation responses.
- gp160 proliferation responses were apparent in 7/32 donors tested, but weaker overall, with a median value for the suppressed group not above that found for HIV seronegative controls
- No differences in the frequency of HIV-specific CD4+ T-cells that were positive for cytokine secretion in a flow cytometry assay were found in the HAART suppressed group versus the group with active viral replication.

HXB2 Location Env
Author Location gp120 (IIIB)
Epitope
Immunogen HIV-1 infection
Species (MHC) human
References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 4/15 responders recognized this peptide, average SI = 4.4.

HXB2 Location Env Author Location gp120 (IIIB) Epitope Immunogen HIV-1 infection Species (MHC) human

References Geretti et al. 1994

- Th proliferative responses were studied in 36 asymptomatic HIV-1 + patients. PBMC from 15/36 patients responded to stimulation with gp120 synthetic overlapping peptide pools. 10 of the responding patients recognized three or more peptide pools.
- After 12 months, most responses were lost or diminished, and specific responses fluctuated. 4/10 of the responders at baseline had new responses at 12 months, as did 5/15 non-responders.
- IL-2 production was a more sensitive and well-preserved measure of Th function than proliferation.
- 4/15 responders recognized this peptide, average SI = 4.4.

HXB2 Location Env
Author Location gp120 (SF2)
Epitope
Subtype B
Immunogen HIV-1 infection
Species (MHC) human

**Keywords** subtype comparisons, rate of progression **References** Imami *et al.* 2002b

• 70 patients with chronic disease progression, 10 clinical nonprogressors, and 3 immunologically discordant progressors (individuals who controlled viremia but had progressive CD4+ T-cell decline) were analyzed for their T-helper cell responses to p24 and cytokine profile.

• In a comparison of responses to HIV-1 proteins based on 10 non-progressors, 3 immunologically discordant, and 70 progressors, SIs were always much higher for non-progressors and immunologically discordant than progressors. Among the non-progressors, the responses to different antigens were greater using p24 peptides than native p24. Native p24, Nef, gp120 proteins, and Remune (gp120 depleted HIV-1, p24 is subtype G), had roughly comparable distributions of SI values from the non-progressors, Nef and gp120 responses were somewhat diminished in immunologically discordant patients.

HXB2 Location Env Author Location (BRU) Epitope Subtype B Immunogen vaccine Vector/Type: inactivated HIV Strain: B clade BRU HIV component: virus Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse

References Haas et al. 1991

- Of 5 mouse inbred lines tested: DBA/2 (H-2d, Ad, Ed), B10.A(4R) (H-2h4, Ak) and B10.A(5R) (H-2i5) showed particularly good CD4+ T cell proliferative responses to HIV proteins (gp160, gp120, p17, p24, Nef and RT), after vaccination with inactivated virus.
- B10.BR (H-2k, Ak, Ek) and C57BL/6 (H-2b and Ab) had weaker responses.

**HXB2 Location** Env

Author Location gp120 (HIV-1,IIIB)

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC) human

Assay type Cytokine production

**Keywords** HIV exposed persistently seronegative (HEPS)

References Fowke et al. 2000

- A cohort of Nairobi sex-worders were defined to be resistant to infection by virtue of remaining seronegative despite repeated high risk exposure. 24 were tested for HIV specific T-helper responses determined by IL-2 production *in vitro* in response to gp120 peptides or soluble gp120 protein.
- In 7/17 resistant women showed IL-2 stimulation >= 2.0, and specific CTL responses were detected in 15/22 resistant women.
   0/12 of the control low-risk subjects had detecatble T-cell responses.

**HXB2 Location** Env

**Author Location** gp160

**Epitope** 

Immunogen HIV-1 infection, vaccine

*Vector/Type:* protein *HIV component:* gp160 *Adjuvant:* aluminum phosphate

Species (MHC) human

Assay type proliferation

Keywords HAART, ART, immunotherapy

References Hejdeman et al. 2003

- Groups of ten asymptomatic HAART-treated HIV-1 + patients with undetected viral loads were monitored for two years after i) no immunization, ii) immunization with rgp160, or iii) immunization with tetanus. Ten HIV-1- volunteers were immunized with tetanus as a control. Results were compared with an rgp160 group tested before HAART was available. The HAART-treated group had increased magnitude and duration of proliferative response to rgp160, maintaining the response for the two year study period. CD4 T-cell responses to tetenus were also improved in the HAART group.
- The recall response to tetanus toxoid and tuberculin were boosted by the rgp160 immunization, particularly in the HAART-treated group.

HXB2 Location Env Author Location

**Epitope** 

Immunogen HIV-1 exposed seronegative

Species (MHC)

Assay type Cytokine production

**Keywords** HIV exposed persistently seronegative (HEPS), acute/early infection, early treatment

References Puro et al. 2000

 This was a case report of a health care worker who had an percutaneous injury and exposure to HIV, and was immediatedly given combination therapy. The individual remained HIV Ab negative, but had transiently detectable viral RNA 2-3 weeks after the exposure. 58 weeks after exposure a Th response was detected by IL-2 production in response to HIV Env peptides.

HXB2 Location Env Author Location Epitope

Immunogen vaccine

Vector/Type: fowlpoxvirus, DNA prime with virus-like particle (VLP) boost Strain: B clade 89.6 HIV component: Env, Gag-Pol

Species (MHC) rabbit

Assay type Cytokine production

Keywords Th1, Th2

References Radaelli et al. 2003

- Rabbits were immunized with fowlpox recombinant vectors or expression plasmids, which express either SIVmac239 gag/pol or HIV-1 env 89.6P genes, and then boosted with virus-like particles (VLPs)(gag/pol SIV with HIV env 89.6).
- A lymphoproliferative Th0 profile response and homologous neutralizing Ab were seen in all three groups. The pcDNA3gag/pol SIV construct was more efficient at producing Abs than the fowlpox construct, although the fowlpox env89.6 construct elicited good humoral and cellular responses. VLP boosting was shown to be efficacious; the pseudoviral structure of the VLP providing a more natural protein conformational was considered helpful for eliciting long term memory cells.

**HXB2 Location** Env **Author Location** gp160

Epitope Subtype B

Immunogen HIV-1 infection, vaccine

Vector/Type: canarypox prime with gp160 boost Strain: B clade MN/LAI-2 HIV component: gp160

Species (MHC) human

Assay type proliferation

**Keywords** vaccine-specific epitope characteristics, vaccine-induced epitopes

References Ratto-Kim et al. 2003

- The CD4+ T-helper response to vaccinees given ALVAC-HIV(vCP205) alone, rgp160 MN/LAI-2 alone, or the two combined in a prime-boost was investigated by establishing T cell lines and comparing proliferative responses to a series of peptides (15 mers overlapping by 10) spanning autologous gp160 MN/LAI-2. Th responses against Env during natural HIV-1 infection were also studied.
- Broad, strong T-helper responses scattered across the Env were obtained from volunteers who received a prime boost vCP205 + rgp160MN/LAI-2, while those receiving rgp160 responded to fewer peptides, and vCP205 to very few peptides.

- HIV-1 + volunteers had less breadth and amplitude of Th responses than vaccinees that got the prime-boost vaccine, although T-cell lines were readily generated from HIV+ individuals. Some vaccinees targeted C1 and C5, while infected individuals did not, and some infected individuals targeted V3, while vaccinees did not.
- The authors note that the differences in response may be contributed to by the fact the peptides used to screen the responses were the same as the vaccine strain, and different than the strains in the natural infection, but that there also may be real immunological differences in the two scenarios of vaccine verses natural infection.

**HXB2 Location** Env **Author Location** gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC)

Assay type proliferation Keywords HAART, ART References Sullivan *et al.* 2003

 Lymphoproliferative responses to HIV antigens p24, gp120 and Nef were enhanced in eight patients who were switched from protease inhibitors to non-nucleoside reverse transcriptase inhibitors.

HXB2 Location Env

 $\textbf{Author Location} \ gp120$ 

**Epitope** 

**Immunogen** HIV-1 infection

Species (MHC) human

Assay type Cytokine production, proliferation

**Keywords** HAART, ART **References** Hardy *et al.* 2003

 Upon initiating HAART, CD4+ T cell proliferative responses in 36 patients were restored to specific antigens, mitogens, and IL-2. Restored recall responses were largely to persistent antigens, and not to HIV-1 or new antigens, and recall responses were associated with IL-2, not IL-4 production.

HXB2 Location Env

Author Location gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords review, Th1, Th2, immune dysfunction

References Becker 2004b

The review suggests HIV-1 shed gp120 virions can act as an allergen, inducing Th2 cytokine production, in particular IL-4, by Fc epsilon RI+ hematopoietic cells. This could inhibit IgG production and CTL responses, and inactivate Th1 cells. New vaccination strategies employing IL-4 inhibitors and antiallergen drugs are discussed.

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords review, Th1, Th2, immune dysfunction

References Becker 2004a

• Review raises the possibility that the switch from Th1 to Th2 activity along with an increase in IL-4 and IgE production in HIV-1 infected patients are an allergic response to HIV-1 protein gp120. Alternative treatments to block Th2 cytokine production, e.g with Il-4 receptor inhibitors, are discussed.

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope Subtype B

Immunogen vaccine

Vector/Type: peptide, heat-shock protein (HSP70) Strain: B clade IIIB HIV component: gp120

Species (MHC) macaque

**Assay type** Cytokine production, proliferation, CD4 T-cell Elispot - IFNγ, T-cell Elispot

**Keywords** genital and mucosal immunity, vaccine antigen design

References Bogers et al. 2004

- Macaques were given vaginal or iliac lymph node immunizations with a novel peptide vaccine composed of SIV p27, CCR5, and N-terminal gp120 fragment, and hsp70 as a carrier.
- 5/8 SHIV 89.6P challenged macaques were protected from infection and vaccinated animals had higher CD4+ T cell numbers than non-vaccinated controls. T-cell proliferation in responses to gp120, vaginal IgG and IgA Abs, and cells producing IL-2, IL-4, and IFNgamma were increased in vaccinated animals.

**HXB2 Location** Env

Author Location gp160 (IIIB)

**Epitope** 

Subtype B

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA, protein, baculovirus Strain: B clade IIIB HIV component: gp160, Nef, Rev, Tat Adjuvant: aluminum phosphate

Species (MHC) human

Country Sweden.

**Assay type** proliferation, CD8 T-cell Elispot - IFNγ

**Keywords** HAART, ART, subtype comparisons, supervised treatment interruptions (STI), immunotherapy

References Boström et al. 2004

- In this study, HIV-infected patients who had previously been immunized with DNA plasmid (nef, tat and ref) or recombinant gp160 were followed longitudinally to determine the impact of HAART on specific T-cell responses. While therapeutic immunizations had transient effects on CD4 cell counts, there was increased survival at 2 years.
- After gp160 vaccination, gp160-specific proliferative CD4+ T cell responses to both baculovirus (MGS HIV-1 rgp 160) and to IMMUNO AG derived gp160 were increased, as well as to p24. Long term HAART treatment was associated with increased IFNγ producing T-cells.
- T-cell proliferative responses to gp160 vaccination were maintained for up to 7 years.

**HXB2 Location** Env

Author Location Env (HXB2. BaL)

Epitope Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade 1007 HIV component: Env, Gag-Pol, Nef

Species (MHC) macaque

**Assay type** CD4 T-cell Elispot - IFNγ, T-cell Elispot **Keywords** variant cross-recognition or crossneutralization, co-receptor

References Letvin et al. 2004

- SIVmac239 gag-pol-nef vaccination of macaques confers better protective responses against a SHIV 89.6 challenge if Env is included even when the vaccine and challenge strain were heterologous in Env. This protection, realized by decreased viral replication and higher levels of CD4+ T cells over time, was associated with T-cell responses early in infection, but not neutralizing Abs.
- The 24 Indian-origin rhesus macaques included in this study did not express Mamu-A\*01.

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype CRF02\_AG

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human

Country Senegal.

Assay type CD4 T-cell Elispot - IFN $\gamma$ 

**Keywords** rate of progression, variant cross-recognition or cross-neutralization

References Zheng et al. 2004

- Gag, Env, Tat, and Nef-specific T-cell responses were evaluated in 68 HIV-1 and 55 HIV-2 infected drug naive, generally asymptomatic, infected Senegalese patients.
- HIV-1 peptides were derived from HIV-1 CRF-02 (HIV-1 A/G, AJ251056)and HIV-2 peptides spanning HIV-2 ROD (M15390).
- Gag specific responses dominated in both groups, but overall
  magnitude and frequencies did not correlate with viral load or
  CD4 counts. More Nef responses were found in HIV-1 infected
  people than HIV-2, and Nef in HIV-2 is more diverse.

## III-B-16 Nef Helper/CD4+ T-cell epitopes

HXB2 Location Nef (1-20)

Author Location Nef (1–20 LAI)

Epitope MGGKWSKSSVVGWPTVRERM

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.

 Proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Nef (1-20)

**Author Location** Nef (1–20 HXB2)

Epitope MGGKWSKSSVIGWPTVRERM

Subtype B

Immunogen HIV-1 infection

Species (MHC) (H-2<sup>d</sup>)

**Keywords** class I down-regulation by Nef **References** Peng & Robert-Guroff 2001

Deletion of the 19 N-terminal amino acids from Nef including the myristolation signal eliminates Nef-induced down-regulation of MHC class I and CD4 molecules. Such a construct has the potential to serve as a more potent immunogen. The known T-cell epitopes that that would be disputed by this deletion are minimal, a murine H-2d Th epitope in the peptide MGGKWSKSSVIGWPTVRERM, and a HLA-B8 CTL epitope, WPTVRERM.

HXB2 Location Nef (14-22)

**Author Location** Nef (14–22 SF2)

**Epitope** SAIRERMRR **Epitope name** 95.12, 33.6

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DRw6)

Donor MHC DRw52, DRw6, DRw15(2), DQw1, DQw6, •

DP4

Assay type proliferation

Keywords epitope processing

References Wentworth & Steimer 1994

- Seven CD4+ T-cell clones that proliferated in responses to SF2
  Nef were generated *in vitro* by stimulation of PBMC from
  uninfected donors. These CD4+ clones also had cytotoxic
  activity.
- These seven clones were capable as of presenting their epitopes to themselves, if Nef peptides were provided.
- The two clones that recognized the epitope SAIRERMRR could also auto-present Nef protein, suggesting that they recognized this epitope in the context of the intact, unprocessed protein.

HXB2 Location Nef (16–35)

**Author Location** Nef (16–35 LAI)

Epitope VRERMRRAEPAADGVGAASR

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

**HXB2 Location** Nef (31–50) **Author Location** Nef (31–50 LAI) Epitope GAASRDLEKHGAITSSNTAA

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Nef (37–51)

**Author Location** 

**Epitope** LEKHGAITSSNTAAT

**Epitope name** N010

Immunogen HIV-1 infection

Species (MHC) human Country Canada.

**Assay type** proliferation, Flow cytometric T-cell cytokine

assay

Keywords memory cells

**References** Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-γ, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2-producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

HXB2 Location Nef (37–51)

**Author Location** 

**Epitope** LEKHGAITSSNTAAT

Epitope name N010

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine assay

Keywords memory cells

References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-γ, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2-producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

HXB2 Location Nef (43-49)

**Author Location** Nef (47–53 SF2)

**Epitope ITSSNTA** 

Epitope name 1.13

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DQw7)

Donor MHC DR1, DR8, DRw52, DQw1, DQw7, DP4

Assay type proliferation

Keywords epitope processing

References Wentworth & Steimer 1994

- Seven CD4+ T-cell clones that proliferated in responses to SF2
  Nef were generated *in vitro* by stimulation of PBMC from
  uninfected donors. These CD4+ clones also had cytotoxic
  activity.
- These seven clones were capable as of presenting their epitopes to themselves, if Nef peptides were provided.

HXB2 Location Nef (45-59)

**Author Location** 

**Epitope** SSNTAATNAACAWLE

Epitope name N012

Immunogen HIV-1 infection

Species (MHC) human Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine

assay

**Keywords** memory cells **References** Younes *et al.* 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-γ, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2-producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

HXB2 Location Nef (45–69)

**Author Location** Nef (45–69 BRU)

Epitope SSNTAATNAACAWLEAQEEEEVGFP

Immunogen vaccine

Vector/Type: peptide prime with protein boost Strain: B clade BRU HIV component: Nef

Species (MHC) chimpanzee, rat

References Estaquier et al. 1992

Antigenic domain: ATNAACAWL, priming with peptide enhanced subsequent Ab response to Nef protein immunization.

HXB2 Location Nef (45-69)

**Author Location** Nef (45–69)

Epitope SSNTAATNAACAWLEAQEEEEVGFP

Immunogen vaccine

Vector/Type: peptide Adjuvant: aluminum

hydroxide

Species (MHC) rat

Keywords vaccine-specific epitope characteristics

References Rouaix et al. 1994

• Covalently linking the potent Th epitope Nef 45-69, which can induce Th proliferative responses at low doses with no adjuvant in Lou/M rats, to a weaker epitope from Schistosoma mansoni allows the induction of detectable Th responses to the Schistosoma epitope.

HXB2 Location Nef (46–65)

**Author Location** Nef (46–65 LAI)

Epitope SNTAATNAACAWLEAQEEEE

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Nef (56-68)

**Author Location** Nef (56–68 HXB2)

Epitope AWLEAQEEEEVGF

Immunogen vaccine

Vector/Type: peptide HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse (DQ2, DQ3, DQ5, DQ6, DQ7, DQ8)

Keywords binding affinity, cross-presentation by differ-

ent HLA, Th1, TCR usage **References** Pancré *et al.* 2002

 This highly conserved Nef epitope has promiscuous HLA-DQ class II binding potential. It has a can bind to 6 different HLA-DQ alleles, but did not bind to any HLA-DR alleles tested. It bound to DQ2 and DQ8 with particularly high affinity, and

with DQ7 with low affinity.

 DQ transgenic mice (in particular DQ8) mounted strong cellular and humoral responses after immunization with this peptide.

- Ex vivo stimulation of CD4+ T-cells from 14 healthy donors (with diverse HLAs) with this peptide presented on autologous DCs resulted in Th1-associated cytokine production. IFNgamma production was stimulated in 7/14 cases, both IFNgamma and IL-2 in 6/14, and just IL-2 in 1/14. No IL-4 or IL-5 production was observed.
- Peptide-specific CD4+ T-cell clones with different HLA presenting molecules demonstrated a preference for TCR V $\beta$ 6.1.

HXB2 Location Nef (61–80)

**Author Location** Nef (61–80 LAI)

Epitope QEEEEVGFPVTPQVPLRPMT

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Hinkula et al. 1997

Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.

• Some proliferative response to vaccination was observed to • These seven clones were capable as of presenting their epitopes peptides throughout Nef and Tat, less for Rev.

HXB2 Location Nef (64-73) **Author Location** Nef (68–77 SF2) Epitope EEVGFPVRPQ

Epitope name 59.25 Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DRw15(2))

Donor MHC DR1, DRw15(2), DQw1, DP4

Assay type proliferation Keywords epitope processing

References Wentworth & Steimer 1994

- Seven CD4+ T-cell clones that proliferated in responses to SF2 Nef were generated in vitro by stimulation of PBMC from uninfected donors. These CD4+ clones also had cytotoxic activity.
- These seven clones were capable as of presenting their epitopes to themselves, if Nef peptides were provided.

HXB2 Location Nef (65-79)

**Author Location** 

Epitope EVGFPVIPQVPLRPM

Epitope name N017

Immunogen HIV-1 infection

Species (MHC) human Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine

assay

Keywords memory cells References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-y, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

HXB2 Location Nef (66-73)

Author Location Nef (70–77 SF2)

Epitope VGFPVRPQ

Epitope name 29.16

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DR1, DRw15(2))

Donor MHC DR1, DRw15(2), DQw1, DP4

Assay type proliferation

Keywords epitope processing

References Wentworth & Steimer 1994

• Seven CD4+ T-cell clones that proliferated in responses to SF2 Nef were generated in vitro by stimulation of PBMC from uninfected donors. These CD4+ clones also had cytotoxic activity.

to themselves, if Nef peptides were provided.

HXB2 Location Nef (66–97)

**Author Location** Nef (66–97 LAI)

Epitope VGFPVTPQVPLRPMTYKAAVDLSHFLKEKGGL

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I trial.
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees – 5/10 reacted to this Nef peptide.
- 9/12 tested mounted a CTL responses to at least one of the six peptides, each of the six peptides elicited a CTL response in at least one individual.
- 5/12 tested had an IgG response to this peptide.

HXB2 Location Nef (76–95)

**Author Location** Nef (76–95 LAI)

Epitope LRPMTYKAAVDLSHFLKEKG

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Nef (81–97)

Author Location Nef (81–97 B Consensus)

Epitope YKAAVDLSHFLKEKGGL

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFN $\gamma$  EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.

• The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

HXB2 Location Nef (91-110) **Author Location** Nef (91–110 LAI)

Epitope LKEKGGLEGLIHSQRRQDIL

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Hinkula et al. 1997

- · Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Nef (98-112)

Author Location Nef (98-112 BRU)

Epitope EGLIHSQRRQDILDL

Immunogen vaccine

*Vector/Type:* peptide prime with protein boost

Strain: B clade BRU HIV component: Nef

Species (MHC) chimpanzee

References Estaquier et al. 1992

• Peptide alone could stimulate monkey T-cells in the absence of carrier protein - required carrier protein in rat.

**HXB2 Location** Nef (104–121)

**Author Location** Nef (104–121 B Consensus)

Epitope QKRQDILDLWVYHTQGYF

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.

• The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** Nef (104–123)

Author Location Nef (106-125 HXB3)

Epitope QRRQDILDLWIYHTQGYFPD?

Immunogen vaccine

Vector/Type: DNA Strain: B clade HXB3

HIV component: Nef

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Sandberg et al. 2000

- · A strong T helper proliferative response against a rec Nef protein was observed 2 weeks after immunization of HLA-A201 transgenic mice in a C57Bl/6 background – the response was weak by 4 weeks post immunization.
- Mice were immunized with nef DNA under the control of a CMV promotor, coated on gold particles delivered to abdominal skin by a gene gun.
- Primary responses were directed at peptides 106-125, 166-185, and 181-205, indicating a response to multiple epitopes.

**HXB2 Location** Nef (106–125)

**Author Location** Nef (106–125 LAI)

Epitope RQDILDLWIYHTQGYFPDWQ

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

Species (MHC) mouse (H-2<sup>b</sup>)

References Hinkula et al. 1997

- · Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Nef (112-127)

**Author Location** Nef (112–127 B Consensus)

Epitope LWVYHTQGYFPDWQNY

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFNγ, Intracellular cytokine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance, acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 14% of the study group.

- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** Nef (112–128) **Author Location** Nef (111–128)

Epitope LWVYHTGQGYFPDWQNYT

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production

References Geels et al. 2006

- The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.
- Autologous sequences corresponding to known and predicted The pitopes were analyzed. LWVYHTGQGYFPDWQNYT had fixation of 1 mutation (LWVYHTGQGYFPDW[q/d]NYT) in 1 of the patients.

**HXB2 Location** Nef (117–147)

**Author Location** Nef (117–147 LAI)

Epitope TQGYFPDWQNYTPGPGVRYPLTFGWCYKLVP

Subtype B

Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I trial.
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees 1/10 reacted to this Nef peptide.
- 9/12 tested mounted a CTL responses to at least one of the six peptides, each of the six peptides elicited a CTL response in at least one individual.
- 10/12 tested had an IgG response to this peptide.

**HXB2 Location** Nef (121–140)

**Author Location** Nef (121–140 LAI)

Epitope FPDWQNYTPGPGVRYPLTFG

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

**HXB2 Location** Nef (136–155)

**Author Location** Nef (136–155 LAI)

Epitope PLTFGWCYKLVPVEPDKVEE

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

**HXB2 Location** Nef (151–170)

**Author Location** Nef (151–170 LAI)

**Epitope** DKVEEANKGENTSLLHPVSL

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

HXB2 Location Nef (162-178)

Author Location Nef (162–178 B Consensus)

Epitope NSLLHPMSLHGMDDPEK

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 11% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.

• The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** Nef (164–183)

Author Location Nef (166–185 HXB3)

Epitope LLHPVSLHGMDDPEREVLEW?

Immunogen vaccine

Vector/Type: DNA Strain: B clade HXB3

HIV component: Nef

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Sandberg et al. 2000

- A strong T helper proliferative response against a rec Nef protein was observed 2 weeks after immunization of HLA-A201 transgenic mice in a C57Bl/6 background – the response was weak by 4 weeks post immunization.
- Mice were immunized with nef DNA under the control of a CMV promotor, coated on gold particles delivered to abdominal skin by a gene gun.
- Primary responses were directed at peptides 106-125, 166-185, and 181-205, indicating a response to multiple epitopes.

**HXB2 Location** Nef (166–185)

Author Location Nef (166–185 LAI)

Epitope HPVSLHGMDDPEREVLEWRF

Subtype B

Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>b</sup>, H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

**HXB2 Location** Nef (176–193)

Author Location Nef (176–193 B consensus)

Epitope PEKEVLVWKFDSRLAFHH

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101, DRB1\*0401,

DRB1\*0701, DRB1\*1101, DRB1\*1302,

DRB1\*1501, DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

Keywords supervised treatment interruptions (STI),

rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFN $\gamma$  EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 36% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The peptides that were recognized by the most people were able to bind broadly to multiple HLA-DR molecules. This peptide showed high cross-reactive binding capacity and bound to 7/8 tested HLA-DR molecules.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** Nef (179–198)

Author Location Nef (181–205 HXB3)

Epitope EVLEWRFDSRLAFHHVAREL?

Immunogen vaccine

Vector/Type: DNA Strain: B clade HXB3

HIV component: Nef

**Species (MHC)** mouse (H-2<sup>b</sup>)

References Sandberg et al. 2000

- A strong T helper proliferative response against a rec Nef protein was observed 2 weeks after immunization of HLA-A201 transgenic mice in a C57Bl/6 background – the response was weak by 4 weeks post immunization.
- Mice were immunized with nef DNA under the control of a CMV promotor, coated on gold particles delivered to abdominal skin by a gene gun.
- Primary responses were directed at peptides 106-125, 166-185, and 181-205, indicating a response to multiple epitopes.

HXB2 Location Nef (181–188)

**Author Location** Nef (185–192 SF2)

Epitope LVWRFDSK

Epitope name 6.38

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DP5)

Donor MHC DRw11, DRw52, DQw7, DP5

Assay type proliferation

**Keywords** epitope processing

References Wentworth & Steimer 1994

- Seven CD4+ T-cell clones that proliferated in responses to SF2
  Nef were generated *in vitro* by stimulation of PBMC from
  uninfected donors. These CD4+ clones also had cytotoxic
  activity.
- These seven clones were capable as of presenting their epitopes to themselves, if Nef peptides were provided.

**HXB2 Location** Nef (181–205) **Author Location** Nef (181–205 LAI) **Epitope** LEWRFDSRLAFHHVARELHPEYFKN

Subtype B Immunogen vaccine

Vector/Type: DNA Strain: B clade LAI

HIV component: Nef, Rev, Tat

**Species (MHC)** mouse (H-2<sup>d</sup>)

References Hinkula et al. 1997

- Stronger, broader responses were observed in animals vaccinated with DNA epidermally rather than with intramuscular protein.
- Some proliferative response to vaccination was observed to peptides throughout Nef and Tat, less for Rev.

**HXB2 Location** Nef (182–205)

**Author Location** Nef (182–205 LAI)

Epitope EWRFDSRLAFHHVARELHPEYFKN

Subtype B Immunogen vaccine

Vector/Type: lipopeptide

Species (MHC) human

References Gahery-Segard et al. 2000

- Anti-HIV lipopeptide vaccine consisting of six long peptides derived from Nef, Gag and Env HIV-1 proteins modified by a palmitoyl chain was administered in a phase I trial.
- A CD4+ T cell proliferative response to at least one of the six peptides was observed in 9/10 vaccinees - 4/10 reacted to this Nef peptide.
- 9/12 tested mounted a CTL responses to at least one of the six peptides, each of the six peptides elicited a CTL response in at least one individual.
- None of the 12 tested had an IgG response to this peptide.

**HXB2 Location** Nef (184–199)

**Author Location** Nef (184–199 B consensus)

Epitope KFDSRLAFHHMARELH

Subtype B

Immunogen HIV-1 infection

Species (MHC) human (DRB1\*0101, DRB1\*0701, DRB1\*1101, DRB1\*1501, DRB5\*0101)

Country United States.

Assay type CD4 T-cell Elispot - IFNγ

**Keywords** supervised treatment interruptions (STI), immunodominance

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNy EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 25% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The peptides that were recognized by the most people were able to bind broadly to multiple HLA-DR molecules. This peptide showed high cross-reactive binding capacity and bound to 5/8 tested HLA-DR molecules.

• The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** Nef (184–199)

Author Location Nef (184–199)

Epitope KFDSRLAFHHMARELH

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country Netherlands.

Assay type Cytokine production

References Geels et al. 2006

- · The relationship between CTL escape and the subsequent increase in viral load and CD4 Th responses was studied in 2 patients. In both patients T-cell reactivity and recognition were lost after CTL escape, and in 1 patient only the loss of CTL responses was paralleled by a decrease in IL-2 CD4 Th responses.
- · Autologous sequences corresponding to known and predicted Th epitopes were analyzed. KFDSRLAFHHMARELH had fixation of 2 mutations (KFDS[r/h]LAF[h/r]HMARELH) in 1 of the patients.

**HXB2 Location** Nef (185–200)

**Author Location** Nef (183–198)

Epitope FDSRLAFHHVARELHP

Immunogen HIV-1 infection

Species (MHC) human

References Ranki et al. 1997

• T-cell response to this epitope persisted after seroreversion.

**HXB2 Location** Nef (186–206)

Author Location Nef (p27) (185–205 BRU)

Epitope DSRLAFHHVARELHPEYFKNC

Epitope name PF63

Subtype B

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU HIV component: gp160, Nef, p17/p24 Gag, p25 Gag Adjuvant: muramyl-dipeptide base

adjuvant (Syntex)

Species (MHC) chimpanzee

Keywords immunodominance

References Bahraoui et al. 1990

- Six chimpanzees were immunized with rec vaccinia viruses (VV) expressing HIV-1 gp160, Gag, and Nef.
- 2/6 chimpanzees showed persistent T-helper proliferative responses against a putative immunodominant epitope located at the C-term end of Nef.

**HXB2 Location** Nef (189–203)

**Author Location** 

**Epitope** LAFHHVARELHPEYF

**Epitope name** N048

Immunogen HIV-1 infection

Species (MHC) human

Country Canada.

Assay type proliferation, Flow cytometric T-cell cytokine

assay

Keywords memory cells

References Younes et al. 2003

- HIV-1-specific CD4+ T-cell responses were analyzed for 6 years since primary infection, in 10 aviremic and 8 viremic patients.
- CD4+ T cells proliferating responses were correlated with the frequency of CD4+ T cells secreting IL-2 in aviremic patients. In viremic patients, CD4+ T cell proliferative response was impaired despite of the high frequencies of IFN-γ, but not IL-2-producing CD4+ T cells in periods of elevated viremia, suggesting that long-term CD4+ memory depends on IL-2-producing CD4+ T cells and that IFN-γ only-producing cells are short lived.

**HXB2 Location** Nef (190–206)

**Author Location** Nef (190–206 B Consensus)

Epitope AFHHMARELHPEYYKDC

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Country United States.

Assay type CD4 T-cell Elispot - IFN $\gamma$ , Intracellular cyto-

kine staining, HLA binding

**Keywords** supervised treatment interruptions (STI), rate of progression, immunodominance,

acute/early infection

References Kaufmann et al. 2004

- CD4 T-cell responses in 36 HIV-1 positive individuals in different disease stages were mapped by IFNγ EliSpot using a panel of 410 B-consensus peptides spanning all HIV proteins. Virus specific CD4+ T-cell responses (median=7, range of magnitude=990) were detected in 30/36 patients.
- This peptide was recognized by 14% of the study group.
- Gag and Nef responses dominated the CD4+ T-cell EliSpot responses in the study. No correlation between plasma viremia and the frequency or magnitude of HIV-1-specific responses was observed.
- The most HIV-1 specific CD4+ T-cell responses were observed in acutely infected patients with treatment of primary infection who stopped after 1-3 treatments (acute STI), compared to patients continuing therapy. People with therapy during chronic infection had low responses; higher numbers of responses were seen among untreated people. 2/9 LTNPs responded to many peptides, comparable to acute STI.

**HXB2 Location** Nef (191–199)

Author Location Nef (195–203 SF2)

Epitope FHHMARELH

Epitope name 3.2

Subtype B

Immunogen in vitro stimulation or selection

Species (MHC) human (DR1)

Donor MHC DR1, DR8, DRw52, DQw1, DQw7, DP4

Assay type proliferation

Keywords epitope processing

References Wentworth & Steimer 1994

- Seven CD4+ T-cell clones that proliferated in responses to SF2
  Nef were generated *in vitro* by stimulation of PBMC from
  uninfected donors. These CD4+ clones also had cytotoxic
  activity.
- These seven clones were capable as of presenting their epitopes to themselves, if Nef peptides were provided.

**HXB2 Location** Nef

**Author Location** Nef

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Nef,

Vif, Vpu

**Species (MHC)** mouse (H-2<sup>d</sup>)

Keywords subtype comparisons, Th1

References Ayyavoo et al. 2000

- Splenocytes from BALB/c mice immunized with pVVN-P DNA were incubated with Vif, Vpu or Nef antigens for 3 days and assayed for IL-4 and IFNγ levels.
- IL-4 production was not significantly changed after antigen stimulation compared to control levels.
- Cross-clade CTL activity was also observed: A, B clade, CRF01(AE) clade antigens could serve as targets for the B clade immunization stimulated CTL – an HIV-1 AC recombinant, however, did not stimulate a CTL response, but was expressed at lower levels on the target cell.

**HXB2 Location** Nef

Author Location Nef (LAI)

**Epitope** 

Subtype B

Immunogen HIV-1 infection

Species (MHC) human

References da Silva & Hughes 1998

- This study compares the level of variation in Nef CTL epitopes to helper and MAb epitopes from the same region.
- CTL epitopes tend to be more conserved than either helper or MAb epitopes and there are stronger functional constraints in the regions where CTL epitopes cluster.

**HXB2 Location** Nef

**Author Location** Nef

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Nef,

Rev, Tat

Species (MHC) human

Keywords HAART, ART

References Calarota et al. 1999

- 9/9 HIV-1 + subjects were given one of three DNA vaccinations for nef, rev or tat, and novel proliferative and CTL responses were generated.
- The nef DNA immunization induced the highest and most consistent CTLp activity, IFN $\gamma$  production, and IL-6 and IgG responses.

 Highly active antiretroviral treatment (HAART) did not induce new HIV-specific CTL responses but reduced viral load, while DNA vaccination induced new immune responses but did not reduce viral load – thus this is a potentially complementary and promising combination.

HXB2 Location Nef Author Location Nef Epitope

Immunogen HIV-1 infection, vaccine

Vector/Type: DNA HIV component: Nef, Rev, Tat Adjuvant: CpG immunostimulatory sequence (ISS)

Species (MHC) human
Keywords review, Th1

References Calarota & Wahren 2001

This review discusses the cellular immune response, and comments on CpG induction of Th1 cytokines and enhanced immune responses, and HIV-1 DNA vaccine boosting of CTL and Th proliferative responses in asymptomatic HIV+ individuals.

HXB2 Location Nef Author Location Nef Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** HAART, ART **References** Oxenius *et al.* 2000

 Patients who started therapy at acute HIV infection (three with sustained therapy, two with limited therapy upon early infection) had strong HIV specific CD4 proliferative responses and were able to maintain a CTL response even with undetectable viral load – three patients that had delayed initiation of HAART had no HIV specific CD4 proliferative responses and lost their CTL responses when HAART was eventually given and their viral loads became undetectable.

HXB2 Location Nef Author Location Nef Epitope

Immunogen HIV-1 infection

Species (MHC) human

Keywords rate of progression, Th1, Th2

References Wilson et al. 2000b

- Dysfunction of HIV-1 specific proliferative responses, but not responses to other antigens, is evident in HIV-1 progressive disease.
- Vigorous HIV-1 specific responses to p24, Nef and gp120 with SI between 8-99 were seen in 6/7 long term non-progressors (LTNP), the seventh had a borderline responses. IL-2 production was seen in all cases, and IL-4 production was also evident many responses.
- None of the progressors (0/5) had HIV-1 specific proliferative responses, or IL-2 or IL-4 induction.
- Non-HIV antigens (cytomegalovirus, PPD, Staphylococcus enterotoxin B, tetanus toxoid) gave similar responses in HIV-1 + LTNP, progressors, and HIV-1 controls.

**HXB2 Location** Nef **Author Location** Nef (BRU)

**Epitope** 

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA), PLG

Species (MHC) mouse

Keywords Th2

References Moureau et al. 2002

- BALB/c mice were immunized with Nef alone, Nef with Freund's adjuvant, or Nef encapsulated in poly(DL-lactideco-glycolide) PLG microparticles.
- High Ab titers (predominantly IgG1) against Nef were retained for seven months in the mice infected with Nef-PLG, 3-fold higher than Nef in Freund's, 5-fold higher than Nef alone.
- CD4+ T-cell lymphoproliferative were observed, and cytokine profiles indicated this was primarily a Th2 response.

**HXB2 Location** Nef **Author Location** Nef (SF2)

Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** subtype comparisons, rate of progression **References** Imami *et al.* 2002b

- 70 patients with chronic disease progression, 10 clinical non-progressors, and 3 immunologically discordant progressors (individuals who controlled viremia but had progressive CD4+T-cell decline) were analyzed for their T-helper cell responses to p24 and cytokine profile.
- In a comparison of responses to HIV-1 proteins based on 10 non-progressors, 3 immunologically discordant, and 70 progressors, SIs were always much higher for non-progressors and immunologically discordant than progressors. Among the non-progressors, the responses to different antigens were greater using p24 peptides than native p24. Native p24, Nef, gp120 proteins, and Remune (gp120 depleted HIV-1, p24 is subtype G), had roughly comparable distributions of SI values from the non-progressors, Nef and gp120 responses were somewhat diminished in immunologically discordant patients.

**HXB2 Location** Nef

**Author Location (BRU)** 

Epitope Subtype B

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade BRU HIV component: RT, virus Adjuvant: Complete Freund's Adjuvant (CFA)

Species (MHC) mouse

References Haas et al. 1991

- Of 5 mouse inbred lines tested: DBA/2 (H-2d, Ad, Ed), B10.A(4R) (H-2h4, Ak) and B10.A(5R) (H-2i5) showed particularly good CD4+ T cell proliferative responses to HIV proteins (gp160, gp120, p17, p24, Nef and RT), after vaccination with inactivated virus.
- B10.BR (H-2k, Ak, Ek) and C57BL/6 (H-2b and Ab) had weaker responses.

HXB2 Location Nef

**Author Location Nef** 

**Epitope** 

Immunogen

Species (MHC)

References

**HXB2 Location** Nef **Author Location Nef Epitope** 

Immunogen HIV-1 infection

Species (MHC)

Assay type proliferation Keywords HAART, ART References Sullivan et al. 2003

• Lymphoproliferative responses to HIV antigens p24, gp120 and Nef were enhanced in eight patients who were switched from protease inhibitors to non-nucleoside reverse transcriptase inhibitors.

**HXB2 Location** Nef **Author Location Nef** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type Cytokine production, proliferation

Keywords HAART, ART References Hardy et al. 2003

• Upon initiating HAART, CD4+ T cell proliferative responses in 36 patients were restored to specific antigens, mitogens, and IL-2. Restored recall responses were largely to persistent antigens, and not to HIV-1 or new antigens, and recall responses were associated with IL-2, not IL-4 production.

#### HIV-1 Helper/CD4+ T-cell III-B-17 epitopes

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** review, HIV exposed persistently seronegative

(HEPS), mother-to-infant transmission

References Kuhn et al. 2002

• Intrauterine exposure of infants to HIV from their mothers results in HIV-1 specific T-helper cell proliferative responses in 1/3 of exposed uninfected babies, and HIV-1 specific CTL in some. It is unknown whether these responses are associated with lack of infection, but there is some evidence that HIV-1 T-cell responses may reduce transmission in breastfeeding mothers. Summary tables are provided of CD4 and CD8 T-cell responses detected in earlier studies.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)) Strain: AG recombinant HZ321 HIV component: gp120 depleted virus Adjuvant: Incomplete Freund's Adiuvant (IFA)

Species (MHC) human

**Keywords** HAART, ART, rate of progression

References Kahn et al. 2000

• No benefit was observed in terms of progression free survival for HIV-1 patients on ART given vaccinations with HIV-1 antigen (N=1,262) versus those vaccinated with placebo (N=1,265). There was no statistically different outcome in HIV RNA, CD4 percentage, or body weight. HIV-1 ART patients that were vaccinated did have higher absolute CD4 counts.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)) Strain: AG recombinant HZ321 HIV component: gp120 depleted virus Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human

Keywords HAART, ART References Moss et al. 1999

• 15 HIV-1 + patients on ARV given vaccinations with HIV-1 antigen versus vaccinated with placebo. Lymphocyte proliferation of CD4+, CD8+ memory cells and NK cells to p24 and Remune HIV-1 antigen increased in HAART treated patients after vaccination.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)) Strain: AG recombinant HZ321 HIV component: gp120 depleted virus Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human

Keywords Th1

References Moss et al. 1997

• HIV-1 specific stimulation of T-cell proliferation, and betachemokines (RANTES) and Th1-type cytokine (IFNgamma) production are found after immunization of HIV-1 + individuals with HIV-1 immunogen.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)) Strain: AG recombinant HZ321 HIV component: gp120 depleted virus Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human

#### References Levine et al. 1996

• Long-term follow up of HIV-1 + individuals given HIV-1 immunogen, suggesting those patients who became HIV-DTH-responsive in response to the HIV-1 immunogen had a better clinical outcome. Of twelve who developed DTH-responsiveness, one got an opportunistic infection and died, and one developed KS. Of the 13 patients who remained HIV-DTH-nonresponsive, 9 (69%) progressed to AIDS and 7 of these had died.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Vector/Type: HIV-1 immunogen Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (MHC) human

References Turner et al. 1994

A dose response study of HIV immunogen in IFA was conducted. Doses of 50, 100, 200, or 400 micrograms (total protein) were tested by DTH skin testing to the inactivated HIV-1 antigen. The HIV-1 immunogen was well tolerated, and the minimum dose required to induce HIV-1 DTH was 100 micrograms.

**HXB2** Location HIV-1

**Author Location** 

Epitope

Immunogen HIV-1 infection Species (MHC) human, macaque

Keywords dynamics, HAART, ART

References Wodarz 2002

 Mathematical modeling is used to support the idea that T-helper cell dysfunction results in a compromised ability to maintain an anti-HIV CTL memory response. Models suggest strategies to restore CTL memory through therapy and improve long-term immunological control of the virus.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

*Vector/Type:* DNA, canarypox, gp120 depleted virus HZ321 (REMUNE(TM)), protein, virus-like particle (VLP), adenovirus *Adjuvant:* GM-CSF, Growth Hormone, IL-12, IL-2, IL-7, CpG immunostimulatory sequence (ISS), Thymosin  $\alpha$ -1

Species (MHC) human

**Keywords** HAART, ART, review, rate of progression, immunotherapy

References Imami et al. 2002a

This review addresses the use of immunotherapy and therapeutic immunization to help chronically infected patients maintain a strong anti-HIV-1 T-cell response. The loss of anti HIV-1 proliferative responses early after infection is reviewed, as are therapeutic vaccinations, with or without HAART, and strategies for immunomodulation that can be given with or without vaccination.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords review, rate of progression, Th1, Th2

**References** Heeney 2002

Review of the importance of balanced Th1 and Th2 HIV-specific CD4 T-cell responses in control of infection and for vaccination strategies.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC)

Keywords dynamics, rate of progression, escape

References Bernaschi & Castiglione 2002

 A cellular automata model was used to model the dynamics of HIV-1 infection and progression to disease. The model suggests the long aymptomatic period is due to immune escape mutants with lower viral fitness, and with AIDS resulting from a drastic reduction of the T-helper cell reservoire.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Species (MHC)

Keywords dynamics, kinetics

References Altes et al. 2002

- This study employs a mathematical model to study the consequences of increasing the T-helper response through a vaccine, which would have counter-balancing effects in a new infection: a more intense response provides more help but also more target cells. The model indicates that if the infecting virus had a low replication rate, then CTLp and CD4 helper cells could control an infection. Only a vaccine that could increase CTL responsiveness could reduce viral set point with observed replication rates.
- A CD4+ T-cell response without maintained CTL response was deleterious in this model.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC)

**Keywords** dynamics, HAART, ART, rate of progression **References** Bajaria *et al.* 2002

 This paper presents a dynamical model of HIV infection and progression that includes CD4 T-cell naive and memory populations distributed between the peripheral blood and the lymph nodes, as well as the effects of HAART. Increasing viral replication and infectivity and decreasing T-cell immunity had impact on the rate of disease progression in this model.

HXB2 Location HIV-1

**Author Location** (HZ321)

**Epitope** 

Subtype AG Immunogen vaccine

Vector/Type: gp120 depleted virus HZ321 (REMUNE(TM)) Strain: AG recombinant HZ321 Adjuvant: Incomplete Freund's Adjuvant (IFA), CpG immunostimulatory sequence (ISS)

Species (MHC) mouse Keywords Th1, Th2

References Ayash-Rashkovsky et al. 2002

• Parasitic helminthic infections in humans, common in parts of Africa and Asia, can shift immune responses to Th2 responses. To model this, BALB/c mice were infected with the parasite Schistosoma mansoni, and the infected mice showed a dominant Th2 immune response. Vaccination with gp120-depleted HIV-1 viral particles and incomplete Freund's adjuvant induced Th2 responses in these mice, but this could be shifted towards a Th1 profile when CpG oligodeoxynucleotide was added to the vaccine as an immunostimulatory agent.

**HXB2** Location HIV-1

Author Location HIV-1 except gp120

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, rate of progression

References Ghanekar et al. 2001

- 12 long term non-progressors (>10 years) went on HAART, while 14 elected not to go on HAART. After a year on HAART, higher frequencies and absolute numbers of HIV-specific memory CD4+ T-cells were observed in untreated patients than patients receiving HAART therapy, tested by stimulation an proliferation responses to HIV Remune antigen (gp120 depleted vaccine).
- These results indicate a control of viral replication in therapynaive patients may be mediated by their ability to respond to recall viral antigen, and that the diminished response in treated patients may contribute to viral rebound.

HXB2 Location HIV-1 Author Location

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Keywords HAART, ART, Th2

References Pido-Lopez et al. 2002

• The thymic output in HAART-treated HIV-1 infected patients with progressive disease was studied. One patient also receiving steroid treatment therapy had a weak response in a sjTREC assay indicating a dysfunctional thymus, while four patients not on steroids had clear positive sjTREC readings after HAART. Stimulation of PBMC with multiple recall antigens including gp120, p24 and Nef and mitogens, and revealed that in the patient treated with steroids there was and induction of a Th2 type response indicated by increased levels of IL-4 secretion in response to antigen.

HXB2 Location HIV-1 Author Location Epitope Immunogen vaccine

*Vector/Type:* peptide *Adjuvant:* GM-CSF, IL-12, IL-2, IL-4, Tumor Necrosis Factor  $\alpha$  (TNF $\alpha$ )

Species (MHC)

**Assay type** Th support of CTL response

**Keywords** binding affinity, review, Th1, Th2, mucosal immunity

References Berzofsky 2001

Vaccine clusters were constructed containing T helper, CTL aand neutralizing antibody epitopes, and used to immunize mice. Four things were found to enhance the vaccine immune response: i) increasing the affinity of the peptide for the presenting MHC molecule, called epitope enhancement; ii) increasing the avidity of MHC/peptide complex for the T-cell receptor; iii) incorporating cytokines IL-2, GM-CSF, TFN-α, or IL-12 and IL-4 which steer responses towards Th1 or Th2 responses; iv) inducing mucosal immunity specifically, with intrarectal being

HXB2 Location HIV-1

**Author Location** 

most effective.

**Epitope** 

Immunogen vaccine

Vector/Type: DNA HIV component: Env, Gag Adjuvant: B7, GM-CSF, IL-12, IL-15

Species (MHC) human

**Keywords** review, Th1, Th2

References Boyer et al. 2002

The first generation of HIV-1 plasmid vaccines in 167 individuals induced T-helper responses in most vaccine recipients, however CTL responses were below a 20% response rate. REV-independent RNA optimized constructs (pGag and pEnv) as well as B7 costimulatory molecules could significantly enhance CD8 effector cell responses. Co-administered GM-CSF enhanced antibody repsonses, IL-12 CTL production. IL-15 increased T cell expansion without increasing T cell help.

HXB2 Location HIV-1

**Author Location HIV-1** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC)

Assav type Cytokine production

Keywords review

References Breen 2002

• HIV-1 triggers immunological dysfunction in multiple ways, including the loss of CD4-positive T helper cells in quantity and function and hyperactivity and changes in the production and activity of cytokines. The role of pro- and anti-inflamatory cytokines are discussed, including IL-10, which can suppress HIV-1, and IL-1, IL-6, TNFα which up-regulate HIV-1.

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

Assay type Cytokine production, proliferation

References Clerici et al. 1993b

• rCD4-IgG treatment was associated with improved Th cell function measured by IL-2 production in response to alloantigen or PHA, but not to influenza (a recall antigen response), in 9/10 patients. No clinical benefit was evident. rCD40IgG was also shown to block gp120 induced suppression of Th cells *in vitro*. Proposed mechanisms include: inhibiting HIV-cell fusion by blocking the binding of gp120 to CD4, competing with free gp120 for binding to the CD4 receptor and reducing gp120 induced immunosupression, and gp120-induced direct killing of Th cells.

HXB2 Location HIV-1 Author Location Nef Epitope Subtype B

Immunogen HIV-1 infection

Species (MHC) human

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ, Intracellular cytokine staining

**Keywords** assay standardization/improvement

References Draenert et al. 2003

• Six different HIV-1 Nef peptide sets ranging in length from 15-20 amino acids with overlap from 10-11 amino acids derived from either the B clade consensus sequences or the B clade sequence B.AU.AF064676 were used to study the impact of using different peptide design strategies to detect CD4 and CD8 T-cell responses. 20 individuals were tested using the six sets of peptides. 17/20 had CD8 T-cell responses to all peptides sets, and 15 of these 17 had CD4 T-cell responses.

HXB2 Location HIV-1
Author Location Nef
Epitope
Subtype B
Immunogen HIV-1 infection
Species (MHC) human

Assay type CD8 T-cell Elispot - IFNγ, CD4 T-cell Elispot - IFNγ, Intracellular cytokine staining

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- Although there was a trend in detecting more CD8 T cell responses using the shorter 15-mer peptides, longer 20-mers were best for detecting more CD4 T-cell responses, but neither result was statistically significant. Similar results were seen in the 15 to 20 amino acid range for both IFN gamma Elispot and ICS assays.
- Use of the consensus versus the natural strain identified slightly
  increased numbers of reactive peptides. Seven reactive peptides were observed with the B consensus peptides but not the
  B.AU.AF064676 peptides, but on the other hand four reactivites were observed using the B.AU.AF064676 peptides but
  not the consensus.

• Using an overlap of 10 or 11 amino acids did not make a difference.

HXB2 Location HIV-1 Author Location Epitope

Epitope

Immunogen HIV-1 infection

Species (MHC)

**Assay type** Cytokine production **Keywords** HAART, ART **References** Galli *et al.* 2003

- HIV-1-infected women who developed Adepose tissue alterations (ATA) while recieving antiretroviral treatment (ART) had a favorable immunological profile with efficient IL-2 production and T-helper function. The authors suggest that ATA may be related to the ART-driven restoration of immune function.
- The most prominant feature of women with ATA that were recieving ART was increased IL-12 production with a lower TNF alpha and IL-10 synthesis.

HXB2 Location HIV-1
Author Location

**Epitope** 

Immunogen HIV-1 infection

Species (MHC)

Keywords review

References Norris & Rosenberg 2002

This paper reviews the role of Th cells in controlling HIV-1 infection, and in other viral infections. It describes CD4+ T-cell support of Ab production, CTL responses, as well as antiviral cytokine production and infected-cell killing. HIV+ patients with a low viral load and rare vigorous HIV-specific CD4+ proliferative responses, and the benefit of early treatment in preserving Th HIV-specific responses allowing immune control when therapy is subsequently stopped, are described.

HXB2 Location HIV-1

**Author Location** 

Epitope

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** review, rate of progression, acute/early infec-

References Norris & Rosenberg 2001

 This review goes over the evidence for HIV-1 specific Th and CTL responses being critical for inhibiting viral replication. LTNPs and those treated during acute HIV-1 infection generate specific Th responses, but most chronically infected individuals do not.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 and GBV-C co-infection

Species (MHC) human

Assay type Cytokine production

**Keywords** HAART, ART, rate of progression, Th1, Th2

References Nunnari et al. 2003

- HIV-1 positive patients co-infected the GBV-C, the hepatitis G virus, have a longer survival time to AIDs and higher CD4+ T cell counts than patients that were not infected with GBV-C. GBV-C co-infected patients showed an intact Th-1 profile over time, with high serum levels of IL-2 and IL-12, and diminishing Th-2 responses reflected by lower levels of IL-4 and IL-10. The opposite was true for HIV-1 + patients that were not co-infected with GBV-C.
- AIDs progression is slower in patients infected with both HIV-1 and hepatitis G virus. It is unclear whether Th-2 and Th-1 cytokines in co-infected patients show cause or consequence of slower AIDs progression. CD4+ cells may support hepatitis G replication.

**HXB2 Location** HIV-1 **Author Location** HIV-1

**Epitope** 

Immunogen HIV-1 infection

Species (MHC) human

**Keywords** dynamics, acute/early infection **References** Korthals Altes *et al.* 2003

 A model of progression was developed that explicitly assumes CD4+ T-cells are both targets of infection and mediators of the immune response. In this model, high viral inoculum with few initial CD4+ T-cells resulted in target-cell-limited infection and high viral load, but with many CD4+ clones and low initial inoculum, infection was controlled by CD4+ clones.

**HXB2 Location** HIV-1 **Author Location** 

**Epitope** 

Immunogen vaccine

Vector/Type: DNA Adjuvant: GM-CSF, IFN $\gamma$ , IL-12, IL-15, IL-18, IL-1 $\alpha$ , IL-2, IL-2/Ig, MIP-1 $\alpha$ , Tumor Necrosis Factor  $\alpha$  (TNF $\alpha$ ), Tumor Necrosis Factor  $\beta$  (TNF $\beta$ ), M-CSF, G-CSF, IL-8, SDF-1 $\alpha$ , RANTES, MCP1

Species (MHC)

**Keywords** review, Th1, Th2, adjuvant comparison **References** Calarota & Weiner 2004

Review summarizes the developments of DNA vaccine enhancement/modulation by 1) improving Th1 cytokine-encoding plasmids 2) by prime-boost vaccine regimens and 3) by chemokine- or T -cell costimulatroy molecule encoding plasmids. Studies involving many approaches for stimulating Th1 responses upon vaccination are compared, and given the initial promise of these strategies, future studies of coadministration or prime boosting with different combinations are advocated.

HXB2 Location HIV-1

Author Location p24 (HIV-2 ROD, HIV-1 IIIB)

**Epitope** 

Immunogen HIV-1 or HIV-2 infection

Species (MHC) human

Country Gambia.

Assay type Cytokine production, proliferation, CD8 Tcell Elispot - IFNγ, Chromium-release assay

**Keywords** rate of progression **References** Jaye *et al.* 2004

- A comparison of T cell responses in HIV-1 and HIV-2 infected asymptomatic patients with CD4+ cell counts of 20% showed no significant difference between both groups. Viral loads were roughly 20 times greater in HIV-1 positive patients than HIV-2 positive patients.
- 10/20 (50%) of HIV-1 infected patients demonstrated proliferative responses with SI greater than 1.4 to gp120, and 11/20 to p24. 8/29 (29%) of HIV-2 infected patients recognized gp105, and 8/29 (29%) p26. Cytokine responses in both groups did not differ.
- 9/21 (43%) of HIV-1 + and 15/30 (50%) of HIV-2 + patients had cyotoxic T cell responses to Gag, and 3/21 (14%) HIV-1 + and 8/30 (27%) HIV-2 + responded to Pol.

**HXB2** Location HIV-1

**Author Location** 

Epitope

Immunogen HIV-1 infection

Species (MHC) human

Country Spain.

Assay type proliferation, Intracellular cytokine staining

**Keywords** HAART, ART **References** López *et al.* 2004

- A clinical trial compared chronically HIV-1 infected patients who had replaced HAART with didanosine (ddI) and hyroxyurea (HU) were followed for 12 months to an untreated HIV+ group and a group that continued on HAART.
- Approximately 20% of the patients treated with ddI-HU had detectable CD4+ T-cell proliferative responses to Gag and Env in contrast to drug-naive and HAART treated HIV-infected patients, who had few or no responses.
- HIV-specific CD8+ T-cell responses were higher in ddI-HU treated patients than HAART treated patients, even in individuals that maintained undetectable viral loads.

**HXB2 Location** HIV-1

**Author Location** Tat (89.6)

**Epitope** 

Immunogen vaccine

Vector/Type: DNA prime with protein boost, ISCOM Strain: B clade IIIB, SIV HIV component: Env, Gag, Tat Adjuvant: Immune stimulating complexes (ISCOM)

Species (MHC) macaque

Assay type Cytokine production, proliferation, CD8 Tcell Elispot - IFN $\gamma$ 

**Keywords** vaccine-specific epitope characteristics, Th1, Th2, vaccine antigen design

References Mooij et al. 2004

• This study compared vaccinating with Tat alone to vaccinating with Tat+Gag+Env. Rhesus macaques (Macaca mulatta) were intramuscularly immunized with a combination of DNA plasmids (HIV-1 IIIB expressing Tat, SHIV-1 89.6P expressing gp120 and SIV mac239 expressing Gag, followed by three boosts with HIV-1 Tat (IIIB) and Env (89.6, gp140) SIV Gag protein. Animals with multi-antigen vaccination had reduced viremia increased CD4+ T-cell counts.

- Tat-Env-Gag immunized animals had weaker Tat-specific Th responses in comparison to animals immunized with Tat alone; but the response to Tat alone was a Th2 response that did not protect from challenge.
- Immunization with Tat-Env-Gag boosted proliferation of Gagspecific IFN-γ and IL-2 producing cells in 3/4 animals (Th1 and Th2 responses) and induced a Th2-immune response (IL-2, IL-4) to Env.
- CD4+ T helper responses to Tat-Env-Gag immunization were correlated with control and reduction of viremia, suggesting a combination of Th1 and Th2 vaccine responses to multiple HIV antigens is advantageous.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen vaccine

Species (MHC) macaque

Keywords review

References Heeney 2004

- Review discusses the status, design and selection of novel HIV vaccines which elicit strong T-helper responses which can in turn can elicit CTL and Ab responses.
- Review discusses the status, design and selection of novel HIV vaccines which elicit strong T-helper responses which can in turn can elicit CTL and Ab responses.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Immunogen HIV-1 infection, vaccine

Species (MHC) human

Keywords review, immunotherapy, adjuvant comparison

References Wahren & Liu 2004

 This review covers immunotherapeutic vaccines use in combination with antiretrovial therapy and use of vaccination in combination with adjuvants and immunomodulators.

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Immunogen

Species (MHC)

Keywords review, adjuvant comparison

References Mitchison & Sattentau 2005

 Review summarizes mechanisms of immunoregulation relevant for new vaccine development, with a brief summary of adjuvant triggering innate immunity through Toll-like receptors (TLRs), Nod molecules, and other activators. DNA encoded adjuvants that have been tested in DNA vaccines are summarized. The balance between Th1 (CTL activating) and Th2 (B cell activating) responses is discussed, and it is noted that BALB/c mice are predominately Th2 responders, C57BL Th1.

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Subtype C

Immunogen HIV-1 infection

Species (MHC) human

Country South Africa.

Assay type Flow cytometric T-cell cytokine assay

References Ramduth et al. 2005

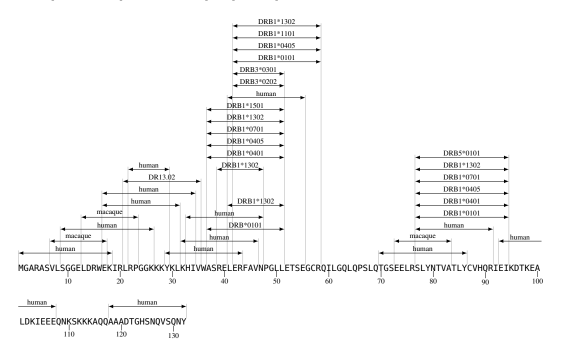
The magnitude of HIV-specific CD8+ T cell responses in HIV-1 infected individuals from South Africa correlated with the CD4+ T cell responses. CD4 responses were narrowly focused, with Gag as dominant target, while CD8 responses were equally distributed among Gag, Pol and the regulatory and accessory proteins. The preferential targeting of Gag by CD8+ T-cells was associated with enhanced control of viral load.

# III-C

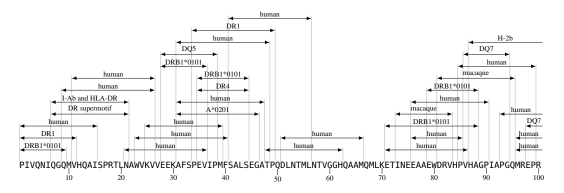
# Maps of T-Helper/CD4 + Epitope Locations Plotted by Protein

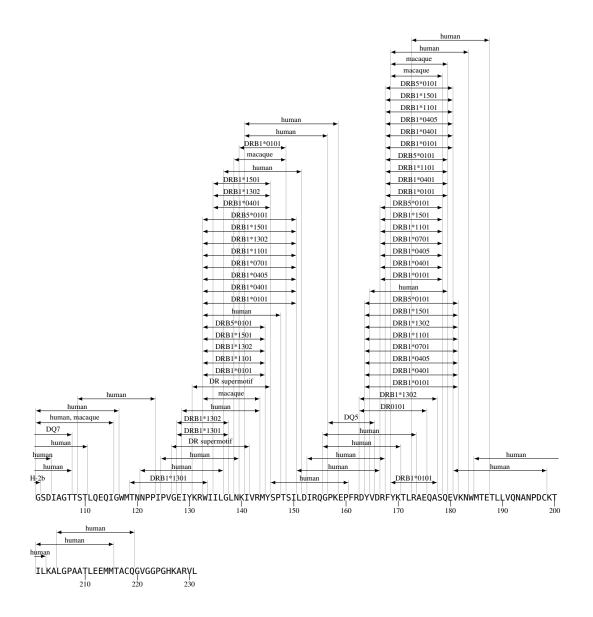
Linear helper T cell (CD4+) epitopes mapped to within a region of 18 amino acids or less are shown.

#### III-C-1 p17 T-Helper/CD4 + Epitope Map

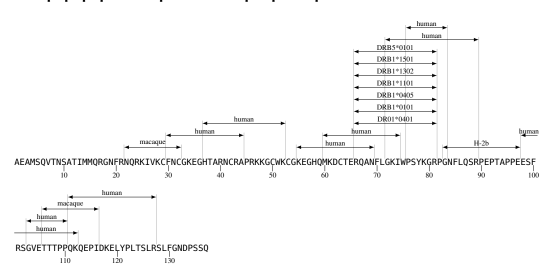


#### III-C-2 p24 T-Helper/CD4 + Epitope Map





#### III-C-3 p2p7p1p6 T-Helper/CD4 + Epitope Map

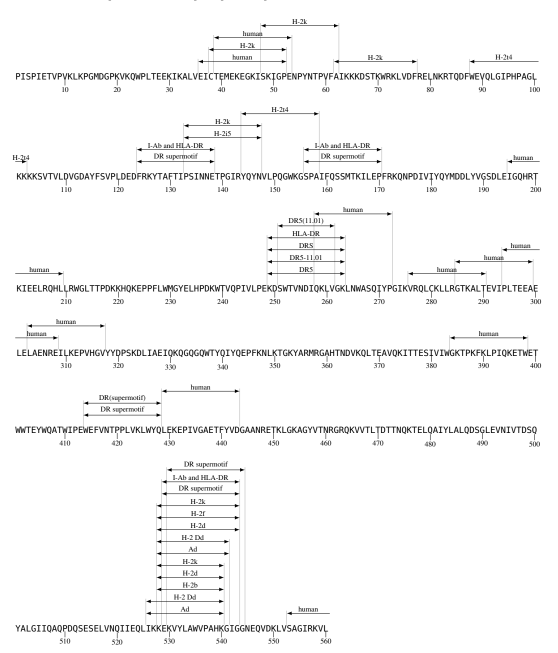


#### III-C-4 Gag/Pol TF T-Helper/CD4+ Epitope Map

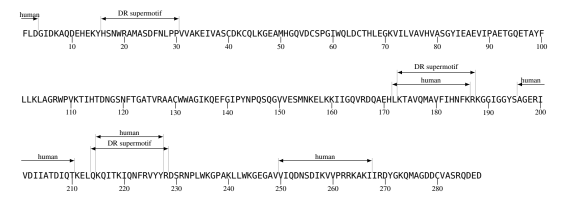
#### III-C-5 Protease T-Helper/CD4 + Epitope Map

 $\verb"PQVTLWQRPLVTIKIGGQLKEALLDTGADDTVLEEMSLPGRWKPKMIGGIGGFIKVRQYPQILIEICGHKAIGTVLVGPTPVNIIGRNLLTQIGCTLNF"$ 

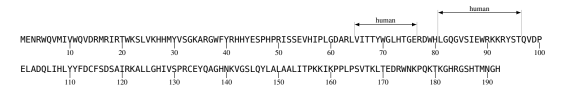
#### III-C-6 RT T-Helper/CD4 + Epitope Map



#### III-C-7 Integrase T-Helper/CD4 + Epitope Map



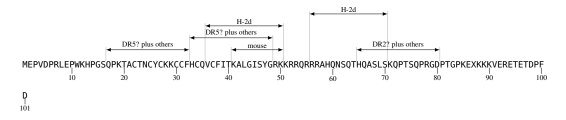
#### III-C-8 Vif T-Helper/CD4 + Epitope Map



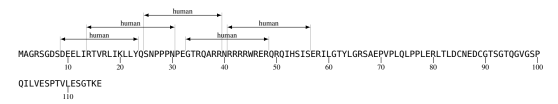
#### III-C-9 Vpr T-Helper/CD4 + Epitope Map



#### III-C-10 Tat T-Helper/CD4+ Epitope Map



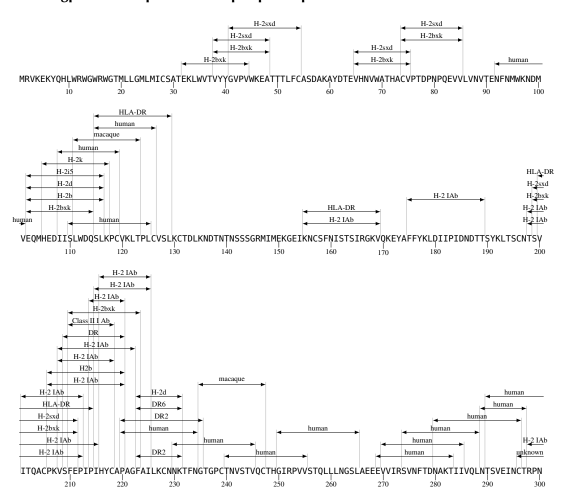
#### III-C-11 Rev T-Helper/CD4 + Epitope Map

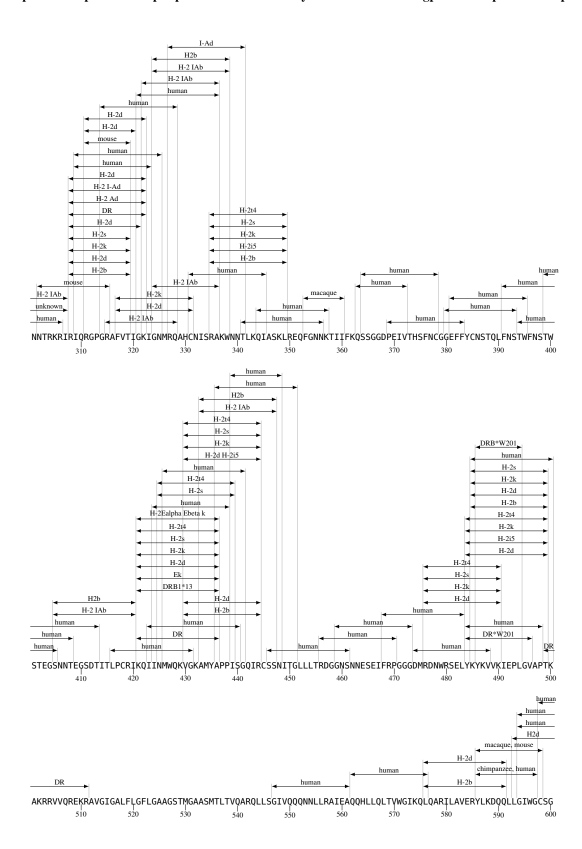


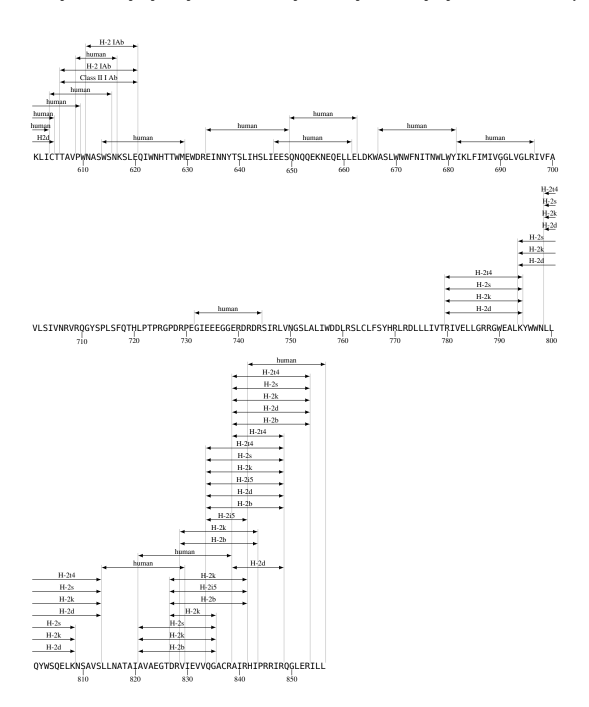
#### III-C-12 Vpu T-Helper/CD4 + Epitope Map



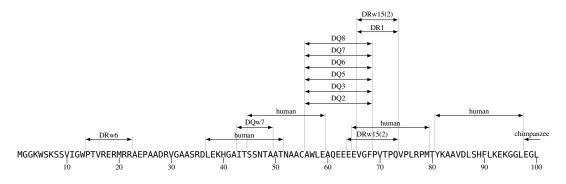
#### III-C-13 gp160 T-Helper/CD4 + Epitope Map

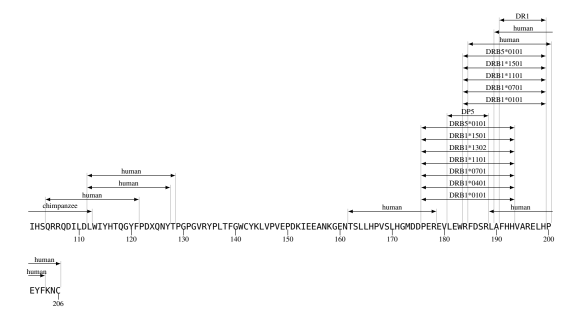






#### III-C-14 Nef T-Helper/CD4+ Epitope Map





# Part IV HIV Antibody Binding Sites

#### IV-A

# **Summary**

This part summarizes HIV-specific antibodies (Abs) arranged sequentially according to the location of their binding domain, organized by protein. We attempted to make this part as comprehensive as possible. For the monoclonal antibodies (MAbs) capable of binding to linear peptides, we require that the binding site be contained within a region of 30 or so amino acids to define the epitope, but not that the precise boundaries be defined. MAbs that do not bind to defined linear peptides are grouped by category at the end of each protein. Antibody categories, for example CD4 binding site (CD4BS) antibodies, are also noted in the index at the beginning of this part. Studies of polyclonal Ab responses are also included. Responses that are just characterized by binding to a protein, with no known specific binding site, are listed at the end of each protein. For more recent updates, epitope sequence alignments, and search capabilities, please see our web site: http: //www.hiv.lanl.gov/content/immunology.

#### **IV-A-1** Indices

Three indices are provided. The first provides a concise list of anti-HIV-1 MAbs by cross-competition category, with both discontinuous epitopes (for example, CD4BS) and some well known linear epitopes (for example, cluster I) summarized. The second lists the MAbs' IDs in alphabetical order so one can find their location in the table. The third is a listing by order of appearance in the tables.

#### **IV-A-2** Tables

Each MAb has a twelve-part basic entry:

**Number:** Order of appearance in this table.

MAb ID: The name of the monoclonal antibody with synonyms in parentheses. MAbs often have several names. For example, punctuation can be lost and names are often shortened (M-70 in one paper can be M70 in another). Polyclonal responses are listed as "polyclonal" in this field.

HXB2 location: Position of the antibody binding site relative to the viral strain HXB2 (GenBank Accession Number K03455), which is used as a reference strain throughout this publication. The numbering in this table corresponds to the protein maps. Because of HIV-1 variation, the epitope may not actually be present in HXB2; rather, the position in HXB2 indicates the position aligned to

the epitope. HXB2 was selected as the reference strain because so many studies use HXB2, and because crystal structures for HXB2-related proteins are available. The precise positions of an epitope on the HXB2 reference strain can be readily obtained using the interactive position locator at our web site: http://www.hiv.lanl.gov/content/hiv-db/LOCATE/locate.html.

Author location: The amino acid positions of the epitope boundaries and the reference sequence used to define the epitope are listed as given in the primary publication. Frequently, these positions as published are imprecise, and do not truly correspond to the numbering of the sequence, but they provide a reasonable guide to the peptide's approximate location in the protein. Also, in many cases, position numbers were provided but the reference sequence identification was not. Because of HIV-1's variability, position numbers require a reference strain to be meaningful. Binding sites that cannot be defined through peptide binding or interference studies are labeled as discontinuous. The approximate location on the protein, sequence number, and reference sequence are listed.

Sequence: The amino acid sequence of the binding region of interest, based on the reference strain used in the study defining the binding site. On occasions when only the position numbers and not the actual peptide sequence were specified in the original publication, we tried to fill in the peptide sequence based on the position numbers and reference strain. If the sequences were numbered inaccurately by the primary authors, or if we made a mistake in this process, we may have misrepresented the binding site's amino acid sequence. Because of this uncertainty, epitopes that were not explicitly written in the primary publication, that we determined by looking up the reference strain and the numbered location, are followed by a question mark in the table.

**Subtype:** The subtype under study, generally not specified for B subtype.

**Neutralizing:** L: neutralizes lab strains. **P**: neutralizes at least some primary isolates. **no**: does not neutralize. No information in this field means that neutralization was either not discussed or unresolved in the primary publications referring to the MAb.

**Immunogen:** The antigenic stimulus of the original B cell response. Often this is an HIV-1 infection. If a vaccine was used as the original antigenic stimulation, not a natural infection, this is noted on a separate line,

and additional information about the vaccine antigen is provided as available.

**Species** (**Isotype**): The host in which the antibody was generated, and the isotype of the antibody.

**Research Contact:** Information about who produced an antibody, how to obtain it, or who should receive credit.

**Country:** The country where the samples were obtained—generally not specified if the study was conducted in the United States.

**References:** All publications that we could find that refer to the use of a specific monoclonal antibody. First is a list of all references. Additional details for some of the older references can be found in Part V, although we have tried to keep the entries self-contained since 1997.

**Keywords:** Keywords for antibody entries were initiated in 2004. The keywords are listed when available as part of the main entry, and also follow the note in bold type so references pertaining to particular types of studies can be found quickly. The keywords include acute/early infection; ADCC; adjuvant comparison; anti-idiotype; antibody binding site definition and exposure; antibody generation; antibody interactions; antibody sequence, variable domain; assay development; assay standardization/improvement; autologous responses; binding affinity; brain/CSF; co-receptor; complement; computational epitope prediction; enhancing activity; escape; genital and mucosal immunity; HAART, ART; HIV exposed persistently seronegative (HEPS); immunodominance; immunoprophylaxis; immunotherapy; immunotoxin; inter-clade comparisons; isotype switch; kinetics; mimics; mimotopes; mother-to-infant transmission; mucosal immunity; neutralization potency; rate of progression; responses in children; reversion, viral fitness; review; structure; subtype comparisons; superinfection; Th1; Th2; vaccine antigen design; vaccine-induced epitopes; vaccine-specific epitope characteristics; and variant cross-recognition or cross-neutralization.

**Notes:** Describe the context of each study, and what was learned about the antibody in the study.

#### IV-A-3 HIV protein binding site maps

The names of MAbs and the location of well characterized linear binding sites of 21 amino acids or less are indicated relative to the protein sequences of the HXB2 clone. This map is meant to provide the relative location of epitopes on a given protein, but the HXB2 sequence may not actually bind to the MAb of interest, as it may vary relative to the sequence for which the epitope was defined. Above each linear binding site, the MAb name is given followed by the species in parentheses. Human is represented by 'h', non-human primate by 'p', mouse by 'm', and others by 'o'. More precise species designations for any given MAb can be found using the web search interface or in the tables in this part.

#### **IV-A-4** Alignments

To conserve space, no epitope alignments are provided in this book, but they can be generated using the Ab search tool at http://www.hiv.lanl.gov/content/immunology. The master alignment files from which the epitope alignments were created are available at our web site at http://www.hiv.lanl.gov/content/hiv-db/ALIGN\_CURRENT/ALIGN-INDEX.html.

# IV-B

# **Cross Reference Listing of MAbs**

#### IV-B-1 MAbs by binding type

Cross reference by protein and binding type of MAb names and their order of appearance in the tables.

Binding type	MAb ID (No.)
p17	
C-term	sc-FV p17 (34)
p24	
C-term	13B5 (116)
Protease	
N-term	1696 (176)
flap region	F11.2.32 (178)
RT	
RT palm domain	6B9 (204)
RT thumb domain	5F (205), 5G (206), 7C4 (207)
gp120 C2	polyclonal (184)
Integrase	
Integrase DNA binding	5D9 (224), 2-19 (227), 8-22 (228), 4-20 (229), 6-19 (230)
domain	
Integrase catalytic core	7-16 (221), 4F6 (222)
N-term	1C4 (208), 2C11 (209), 2E3 (210), 3E11 (211), 3F9 (212), 5F8 (213), 6G5 (214),
	7B6 (215), 7C6 (216), 6C5 (217), 4D6 (220)
gp120 V3	polyclonal (235)
Pol	
C-term	33 (256), F-6 (257)
Vif	
C-term	TG001 (259)
Tat	
C-term	polyclonal (263), polyclonal (274), polyclonal (275), 1D2F11 (277), 2D9E7 (278),
	4B4C4 (279), 5G7D8 (280), NT2/4D5.24 (282), polyclonal (283), 2D9D5 (293),
	polyclonal (294), polyclonal (295), polyclonal (296)
N-term	polyclonal (263), TA9 (265), TD84 (266), TE135 (267), polyclonal (268),
	NT3/2D1.1 (269), 1D9D5 (271), polyclonal (274), polyclonal (275), polyclonal
	(283), polyclonal (294), polyclonal (295), polyclonal (296), G1 (297), G2 (298), J1
	(299), TC15 (300), polyclonal (301), polyclonal (302)
Tat basic region	polyclonal (263), TB12 (272), polyclonal (274), polyclonal (275), polyclonal (276)
	polyclonal (283), polyclonal (294), polyclonal (295), polyclonal (296), B1E3 (303)
	J3B2 (304)
Env (gp160)	
C-HR	polyclonal (994)
C-domain	polyclonal (657), 5B2 (730), 9G11 (731), TH-Ab1 (732), polyclonal (733),
	polyclonal (734), polyclonal (735), polyclonal (736)

Binding type	MAb ID (No.)
C-term	105-306 (627), 750-D (629), 158F3 (632), 161D7 (633), 722-D (635), polyclonal (636), 1131-A (638), 858-D (639), 989-D (640), 14D9 (737), 2F5 (738), 4E10 (739), Z13 (740), C8 (745), 1575 (761), polyclonal (765), polyclonal (766), SAR1 (768), 1577 (769), polyclonal (770), 101-342 (995), 101-451 (996), 120-1 (997), T26 (998), D33 (999), polyclonal (1000)
Env oligomer	T22 (1128)
Leucine zipper motif	(649), (650)
N-HR	polyclonal (994)
N-term	polyclonal (658), D33 (999), 2A2 (1129), AC4 (1130), AD3 (1131), AD3 (1132), ID6 (1133), ID6 (1134)
gp120 C1 gp120 C1-C2	M85 (318), 7E2/4 (319), 4D4#85 (320), M92 (321), M86 (322), polyclonal (323), 133/237 (324), 133/290 (325), 133/11 (326), D/3G5 (327), D/6A11 (328), D/5E12 (329), L5.1 (330), 4A7C6 (331), 1D10 (332), B242 (333), 133/192 (334), 489.1(961) (335), 5B3 (336), B10 (337), B2 (338), C6 (339), MF49.1 (340), T1.1 (341), T7.1 (342), T9 (343), GV4D3 (344), B27 (345), B9 (346), B35 (347), D/4B5 (348), D/5A11 (349), D/6B2 (350), B18 (351), B20 (352), MF39.1 (353), 187.2.1 (354), 37.1.1(ARP 327) (355), 6D8 (356), M96 (357), MF119.1 (358), MF4.1 (359), MF53.1 (360), MF58.1 (361), MF77.1 (362), T2.1 (363), 11/65 (364), W1 (365), T11 (366), GV1A8 (367), 11 (368), 12G10 (369), 135/9 (370), 7C10 (371), C4 (372), MF46.1 (373), 212A (1001), 522-149 (1002), CA1 (1003), CA13 (1004), L19 (1005), M90 (1006), MAG 104 (1007), MAG 45 (1008), MAG 95 (1009), MAG 97 (1010), P35 (1011), T9 (1012), p7 (1013) polyclonal (983), L100 (1014)
gp120 C1-C4	2/11c (1015), A32 (1016)
gp120 C1-C5	C11 (1017), L81 (1018)
gp120 C2	1006-30-D (414), 847-D (415), 213.1 (419), B12 (420), B13 (421), C13 (422), M89 (423), B21 (424), B23 (425), B24 (426), B25 (427), B3 (428), B26 (429), B29 (430), B36 (431), 110.E (432), 110.C (433)
gp120 C3	2H1B (387), 110.D (574), B32 (575), ICR38.1a (587), 2F19C (782), B2C (1019), polyclonal (1020)
gp120 C4	5C2E5 (582), G3-211 (583), G3-537 (584), ICR38.1a (587), G3-299 (588), G3-42 (589), G3-508 (590), G3-519 (591), G3-536 (592), ICR38.8f (593), MO86/C3 (594), 13H8 (595), G45-60 (596), polyclonal (597), 1662 (598), 1663 (599), 1664 (600), 1697 (601), 1794 (602), 1804 (603), 1807 (604), 1808 (605), 1024 (1021), 4KG5 (1022)
gp120 C5	9201 (610), 1C1 (611), 3F5 (612), 5F4/1 (613), 660-178 (614), 9301 (615), B221 (616), H11 (618), W2 (619), M38 (620), 110.1 (623), 42F (624), 43F (625), RV110026 (626), GV1G2 (628), 450-D (630), 670-D (631), 1331A (637), polyclonal (983), 23A (1023), D7324 (1024)
gp120 CCR5BS	E51 (580), 17b (1119), 21c (1120)

Binding type	MAb ID (No.)						
gp120 CD4BS	JL413 (581), polyclonal (585), 1795 (586), polyclonal (983), D33 (999), polyclonal						
-	(1000), 10/46c (1025), 1008-D (1026), 1027-30-D (1027), 1125H (1028), 1125H						
	(1029), 120-1B1 (1030), 1202-D (1031), 1331E (1032), 1570 (1033), 1595 (1034),						
	1599 (1035), 15e (1036), 21h (1037), 28A11/B1 (1038), 2G6 (1039), 35F3/E2						
	(1040), 38G3/A9 (1041), 428 (1042), 448-D (1043), 46D2/D5 (1044), 48-16 (1045),						
	50-61A (1046), 5145A (1047), 558-D (1048), 559/64-D (1049), 55D5/F9 (1050),						
	588-D (1051), 654-D (1052), 67G6/C4 (1053), 729-D (1054), 830D (1055), 9CL						
	(1056), BM12 (1057), D20 (1058), D21 (1059), D24 (1060), D25 (1061), D28						
	(1062), D35 (1063), D39 (1064), D42 (1065), D52 (1066), D53 (1067), D60 (1068),						
	DA48 (1069), DO8i (1070), F105 (1071), F91 (1072), FG39 (1073), Fbb14 (1074),						
	GP13 (1075), GP44 (1076), GP68 (1077), HF1.7 (1078), HT5 (1079), HT6 (1080),						
	HT7 (1081), ICR 39.13g (1082), ICR 39.3b (1083), Ia3 (1084), Ia7 (1085), IgG1b12						
	(1086), IgGCD4 (1087), L28 (1088), L33 (1089), L41 (1090), L42 (1091), L52						
	(1092), L72 (1093), M12 (1094), M13 (1095), M6 (1096), MAG 116 (1097), MAG						
	12B (1098), MAG 29B (1099), MAG 3B (1100), MAG 55 (1101), MAG 72 (1102),						
	MAG 86 (1103), MAG 96 (1104), MTW61D (1105), S1-1 (1106), T13 (1107), T49						
	(1108), T56 (1109), TH9 (1110), anti-CD4BS summary (1111), b11 (1112), b13						
	(1113), b14 (1114), b3 (1115), b6 (1116), polyclonal (1117), (1118)						
gp120 CD4i	D19 (466), E51 (580), (1118), 17b (1119), 21c (1120), 23e (1121), 48d (1122), 49e						
	(1123), Fbb21 (1124), Fbb21 (1125), X5 (1126), 8F101 (1127), 41.1 (1209)						
gp120 V1	35D10/D2 (377), 40H2/C7 (378), 43A3/E4 (379), 43C7/B9 (380), 45D1/B7 (381),						
	46E3/E6 (382), 58E1/B3 (383), 64B9/A6 (384), 69D2/A1 (385), 82D3/C3 (386),						
	polyclonal (606)						
gp120 V1-V2	polyclonal (983), 4KG5 (1022), 11/68b (1135), 62c (1136), CRA-6 (1137), L15						
	(1138), T52 (1139), T54 (1140)						
gp120 V1-V2 and V3-V5	polyclonal (1141)						
gp120 V2	6D5 (374), B33 (375), 697-D (388), C108G (390), 11/4c (395), 8.22.2 (396), 12b						
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	(1143), 1357 (1144), 1361 (1145), 1393A (1146), 2158 (1147), 66a (1148), 66c						
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	SC258 (1155)						
gp120 V2-CD4BS	L25 (1156), L39 (1157), L40 (1158), L78 (1159)						

Binding type	MAb ID (No.)
gp120 V3 gp120 V3 gp120 V3 discontinuous gp120 V3 mimotope gp120 V3-C4 gp120 V3-C5 gp120 V4 gp120 V5 gp120 V5-C5 gp120 adjacent to CD4BS gp120 carbohydrates at glycosylation residues in C2,	(Hilb-V3-26 (434), Hilb-V3-21 (435), 168B8 (436), polyclonal (447), polyclonal (438), MO97/V3 (439), polyclonal (440), 55/11 (441), 8/38c (442), 8/64b (443), polyclonal (444), polyclonal (445), polyclonal (446), polyclonal (447), 9284 (448), polyclonal (444), polyclonal (445), polyclonal (445), polyclonal (445), polyclonal (451), polyclonal (452), MAG 109 (453), MAG 49 (454), MAG 53 (455), MAG 56 (456), 1324-E (457), polyclonal (458), MO99/V3 (459), C311E (460), 924 (462), polyclonal (463), polyclonal (470) cGP 47 439 (471), polyclonal (472), 178.1 (473), 257-D (474), 311-11-D (475), 41148D (476), 391/95-D (477), Aw (478), Bw (479), DO142-10 (480), Dv (481), Fi (482), Gv (483), Hv (484), polyclonal (485), 50.1 (486), (487), BAT123 (488), 838-D (489), 1006-15D (490), 782-D (491), 908-D (492), 1027-15D (493), F19.26-2 (494), F19.48-3 (495), F19.57-11 (496), 13105100 (497), M77 (498), polyclonal (499), SP.BAL114 (500), SP.SF2:104 (501), polyclonal (502), 19b (503), loop 2 (504), 4G10 (505), 5F7 (506), G3-523 (507), MN215 (508), Nea 9301 (509), 4117C (510), 419-D (511), 453-D (512), 504-D (513), 83.1 (514), 5023B (515), F58/D1 (516), P1/D12 (517), P4/D10 (518), IIIB-13 V3 (519), IIIB-34 V3 (520), A47/B1 (521), D59/A2 (522), G44/H7 (523), MO96/V3 (524), μ5.5 (525), 268-D (526), 386-D (527), 5042A (528), 5042B (529), 418-D (530), 5021 (531), 5025B (532), 5042 (533), 110.3 (534), 110.4 (535), 110.5 (536), 58.2 (537), 537-D (539), 5020 (540), RC25 (541), 5023A (542), 110.6 (543), polyclonal (544), 10/36e (545), 10/56 (552), 1034 (553), 591 (554), 901/clonal (548), 0.5β (549), Cβ1, 0.5β (550), NM-01 (551), 1026 (552), 1034 (553), 591 (554), 901/clonal (569), polyclonal (606), 447-52D (744), polyclonal (780), 1334-D (783), C011 (834), F425 B488 (843), polyclonal (983), A6K65 (1022), (1118), 1160), 1108 (1161), 1106 (1162), 110J (1163), 1165 (1164), 2182 (1165), 2191 (1166), 2219 (1167), 2412 (1168), 2442 (1169), 2456 (1170), 2483 (1177), 4919 (1001), 73,C4 (564), MO101/V3,C4 (565), Dolyclonal (1203), polyclonal (1204),
C3, C4, and V4	
gp120-CD4 complex gp41 adjacent to cluster II	8F101 (1127), 8F102 (1242), CG-10 (1243), CG-25 (1244), CG-4 (1245), CG-76 (1246), CG-9 (1247) 14D9 (737), 2F5 (738)
gp41 alpha-helical hairpin intermediate	98-6 (723), polyclonal (1214)

Binding type	MAb ID (No.)
gp41 cluster I	50-69 (662), 246-D (682), 181-D (685), 240-D (686), F240 (687), D49 (688), D61
	(689), T32 (690), T34 (691), 3D6 (718), 1367 (1216), 7B2 (1217)
gp41 cluster II	D50 (720), 98-6 (723), 167-7 (724), ND-15G1 (725), 167-D (726), 126-6 (1218),
	1342 (1219), 1379 (1220), 2.2B (1221), Fab D11 (1222), Fab D5 (1223), Fab G1
	(1224), Fab M10 (1225), Fab M12 (1226), Fab M15 (1227), Fab S10 (1228), Fab S6
	(1229), Fab S8 (1230), Fab S9 (1231), Fab T3 (1232), Md-1 (1233), 1281 (1240)
gp41 cluster III	Fab A9 (1234), Fab G15 (1235), Fab G5 (1236), Fab L1 (1237), Fab L11 (1238), Fab
	L2 (1239)
gp41 cytoplasmic domain	Chessie 8 (1241)
gp41 internal trimeric	1034 (778), 1492 (779)
coiled-coil of N-helices	
gp41 six-helix bundle and the	1010 (772), 1018 (774), 1020 (776), 1022 (777)
internal trimeric coiled-coil of	
N-helices	1(7 D /72() 1014 /772) 1010 /775)1 -11 /004) 1201 /1240) NG 1 /125)
gp41six-helix bundle	167-D (726), 1014 (773), 1019 (775), polyclonal (994), 1281 (1240), NC-1 (1253)
immunodominant region	3D6 (718), 105-518 (1248)
p24+gp41 <b>Nef</b>	31A1 (1249), 39A64 (1250), 39B86 (1251), 9303 (1252)
C-term	AE6 (1298), AG11 (1299), EH1 (1300), AE6 (1308)
HIV-1	ALO (1290), AOTI (1299), EITI (1300), ALO (1300)
RT thumb domain	polyclonal (1359)
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gp120 C1-C4	8.2A (1337), EH21 (1340)
gp120 CCR5BS	1.9E (1314), 1.9F (1315), 2.5E (1327), 4.8E (1334), E047 (1338), ED10 (1339),
	LA15 (1344), LA28 (1346), LF17 (1347)
gp120 CD4BS	13a15 (1318), 13a23 (1319), 13a3 (1320), 13a6 (1321), 13a7 (1322), 13b18 (1323),
	13b53 (1325), 13b61 (1326), 25G (1329), 5145A (1335), 5E (1336), F1 (1341)
gp120 V3	12.19 (1316), 12.9 (1317), 2191 (1328), 2601 (1330), F2A3 (1342), F3.9F (1343),
	LA21 (1345), polyclonal (1361)
gp120 V4	M2 (1348)
gp120 adjacent to CD4BS	1.4C (1312), 1.4G (1313), 4.11C (1332), 4.6H (1333)
quaternary structure	2909 (1331)

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71-31	141	9201	610	B15	577	CA1	1003
714/01	54	9205	566	B18	351	CA13	1003
722-D	635	924	462	B1E3	303	CA13	128
729-D	1054	9284	448	B1E3	338	CB-13/5	42
74	402	9301	615	B20	352	CD-4/1	45
75 75	715	9303	1252	B20 B21	424	CD-4/1 CD12B4	108
750-D	629	9305	1185	B221	616	CD12B4 CD9	148
782-D	491	97B1/E8	822	B23	425	CG-10	1243
7B2	1217	98-4.3	143	B23 B24	426	CG-10 CG-25	1243
7B6	215	98-4.9	144	B242	333	CG-23 CG-4	1244
7C10	371	98-43	664	B25	427	CG-76	1245
7C3	231	98-6	723	B26	429	CG-70 CG-9	1247
7C4	207	989-D	640	B27	345	CG-9 CGP 47 439	471
7C4 7C4	248	9A4C4	105	B27 B29	430	CH9B2	149
7C4 7C6	216	9CL	1056	B2C	1019	Chessie 8	1241
7E2/4	319	9G11	731	B3	428	Chim 1	621
7F11	232	9G11 9G2	310	b3	1115	clone 3	709
7F11	579	9G2 9G5	310	B30	741	CO11	834
/1.11	319	7UJ	31	UCA	/41	COH	034

CRA-3	1152	EH1	1300	FC12	131	ICR 39.3b	1083
CRA-4	1153	EH12E1	151	FF1	74	ICR38.1a	587
CRA-6	1137	EH21	1340	FG39	1073	ICR38.8f	593
CRA1	608	F-6	257	FH2	115	ID6	1133
D/3G5	327	F1	1286	Fv	482	ID6	1134
D/4B5	348	F1	1341	G1	297	ID8F6	40
D/5A11	349	F105	1071	G11G1	152	IE8G2	155
D/5E12	329	F11.2.32	178	G11H3	153	IgA6/30λ	855
D/6A11	328	F14.11	1274	G12	848	IgA6/5k	856
D/6B2	350	F172-D8	719	G2	298	IgA6/L4	857
D/6D1	570	F19.26-4	494	G2 G2	849	IgG1b12	1086
D1	835		495	G3-136	398		
		F19.48-3				IgGCD4	1087
D12	836	F19.57-11	496	G3-1472	1195	IIIB-13 V3	519
D16	837	F2	1279	G3-211	583	IIIB-34 V3	520
D19	466	F223	840	G3-299	588	IIIB-V3-01	569
D20	1058	F240	687	G3-4	399	IIIB-V3-21	435
D21	1059	F285	841	G3-42	589	IIIB-V3-26	434
D24	1060	F2A3	1342	G3-508	590	IVI-4G6	854
D25	1061	F3	1283	G3-519	591	J1	299
D27	1191	F3.9F	1343	G3-523	507	J1	412
D28	1062	F4	1278	G3-536	592	J3	413
D33	999	F424	842	G3-537	584	J3B2	304
D35	1063	F425 B4e8	843	G44/H7	523	J4	260
D39	1064	F5-2	41	G45-60	596	JB7	133
D4	838	F5-4	97	GE4	132	JF11	134
D42	1065	F5.5	1194	GP13	1075	JL413	581
D43	839	F58/D1	516	GP44	1076	K14	858
D47	1192	F7	844	GP68	1077	K24	1196
D49	688	F8	1284	Groo	483	KU32	859
D50	720	F91	1072	GV GV1A8	367	L-anti-Tat	285
D52	1066	Fab A1	667	GV1G2	628	L100	1014
D53	1067	Fab A12	845	GV4D3	344	L14	110
D56	1193	Fab A2	846	GV4H3	411	L14.17	1
D59/A2	522	Fab A4	668	H11	618	L15	1138
D60	1068	Fab A9	1234	H2	850	L17	1154
D61	689	Fab D11	1222	H8	851	L19	1005
D7324	1024	Fab D5	1223	HBW4	852	L25	1156
DA48	1069	Fab G1	1224	HF1.7	1078	L28	1088
DE7	317	Fab G15	1235	HH3	126	L33	1089
DF3	129	Fab G5	1236	HIVIG	853	L39	1157
DG8	124	Fab L1	1237	HT5	1079	L40	1158
DO142-10	480	Fab L11	1238	HT6	1080	L41	1090
DO8i	1070	Fab L2	1239	HT7	1081	L42	1091
Dv	481	Fab L9	847	human sera	158	L5.1	330
DZ	771	Fab M10	1225	Hv	484	L52	1092
E-4	203	Fab M12	1226	HyHIV-1	5	L72	1093
E047	1338	Fab M12B	669	HyHIV-15	24	L78	1159
E5	1290	Fab M15	1227	HyHIV-19	154	L81	1018
E51	580	Fab M26B	670	HyHIV-2	6	LA15	1344
E7	1297	Fab M8B	671	HyHIV-21	15	LA21	1345
E9	1288	Fab S10	1228	HyHIV-22	17	LA28	1346
EB1A9	82	Fab S6	1229	HyHIV-3	7	LA9 (121-134)	760
EB5	125	Fab S8	1230	HyHIV-4	8	LF17	1347
EC3	130	Fab S9	1231	HyHIV-5	9	LH-104-A	89
EC6	122	Fab T2	672	HyHIV-6	10	LH-104-B	118
ED10	1339	Fab T3	1232	i5B11	121	LH-104-C	102
ED6	759	Fbb14	1074	Ia3	1084	LH-104-E	86
ED8	150	Fbb21	1124	Ia7	1085	LH-104-G	120
EF7	85	Fbb21	1125	ICR 39.13g	1082	LH-104-I	119

I II 104 IZ	00	ME97.1	410		160		502
LH-104-K	88 504	MF87.1	418	polyclonal	168	polyclonal	502
loop 2	504	MN215	508	polyclonal	169	polyclonal	538
M-1	693	MO101/V3,C4	563	polyclonal	170	polyclonal	544
M-11	694	MO101/V3,C4	564	polyclonal	171	polyclonal	548
M-13	695	MO101/V3,C4	565	polyclonal	172	polyclonal	555
M-2	696	MO28	862	polyclonal	173	polyclonal	557
M-22	697	MO30	863	polyclonal	174	polyclonal	576
M-24	698	MO43	864	polyclonal	184	polyclonal	585
M-25	699	MO86/C3	594	polyclonal	192	polyclonal	597
M-28	700	MO9.42.2	98	polyclonal	198	polyclonal	606
M-29	701	MO9.50.2	99	polyclonal	235	polyclonal	607
M-36	702	MO96/V3	524	polyclonal	253	polyclonal	622
M-4	703	MO97/V3	439	polyclonal	254	polyclonal	634
M-6	704	MO99/V3	459	polyclonal	255	polyclonal	636
M12	123	MTW61D	1105	polyclonal	261	polyclonal	648
M12	1094	multiple Fabs	876	polyclonal	262	polyclonal	657
M13	1095	multiple MAbs	877	polyclonal	263	polyclonal	658
m18	875	multiple MAbs	878	polyclonal	264	polyclonal	660
M2	1348	multiple MAbs	879	polyclonal	268	polyclonal	673
M25	860	N11-20	558	polyclonal	273	polyclonal	675
M38	620	N2-4	865	polyclonal	274	polyclonal	677
M6	1096	N70-1.9b	560	polyclonal	275	polyclonal	679
M77	498	N70-2.3a	866	polyclonal	276	polyclonal	680
M85	318	NC-1	1253	polyclonal	281	polyclonal	683
M86	322	ND-15G1	725	polyclonal	283	polyclonal	707
M89	423	Nea 9301	509	polyclonal	286	polyclonal	708
M90	1006	NF1A1	1295	polyclonal	287	polyclonal	716
M91	609	NF2B2	1304	polyclonal	288	polyclonal	727
M92	321	NF3A3	1304	polyclonal	289	polyclonal	728
M96	357	NF8B4	1305		290	polyclonal	729
MAb 35	234	NM-01		polyclonal	290		733
			551 282	polyclonal	291	polyclonal	734
MAG 100	1007	NT2/4D5.24		polyclonal		polyclonal	
MAG 109	453	NT3/2D1.1	269	polyclonal	294	polyclonal	735
MAG 116	1097	P1/D12	517	polyclonal	295	polyclonal	736
MAG 12B	1098	P35	1011	polyclonal	296	polyclonal	742
MAG 29B	1099	P4/D10	518	polyclonal	301	polyclonal	765
MAG 3B	1100	P43110	867	polyclonal	302	polyclonal	766
MAG 45	1008	P5-3	868	polyclonal	323	polyclonal	770
MAG 49	454	p7	1013	polyclonal	376	polyclonal	780
MAG 53	455	PC5009	646	polyclonal	405	polyclonal	781
MAG 55	1101	polyclonal	2	polyclonal	437	polyclonal	880
MAG 56	456	polyclonal	23	polyclonal	438	polyclonal	881
MAG 6B	861	polyclonal	36	polyclonal	440	polyclonal	882
MAG 72	1102	polyclonal	43	polyclonal	444	polyclonal	883
MAG 86	1103	polyclonal	48	polyclonal	445	polyclonal	884
MAG 95	1009	polyclonal	55	polyclonal	446	polyclonal	885
MAG 96	1104	polyclonal	80	polyclonal	447	polyclonal	886
MAG 97	1010	polyclonal	83	polyclonal	449	polyclonal	887
Md-1	1233	polyclonal	96	polyclonal	450	polyclonal	888
MF119.1	358	polyclonal	135	polyclonal	451	polyclonal	889
MF169.1	416	polyclonal	159	polyclonal	452	polyclonal	890
MF170.1	417	polyclonal	160	polyclonal	458	polyclonal	891
MF39.1	353	polyclonal	161	polyclonal	463	polyclonal	892
MF4.1	359	polyclonal	162	polyclonal	464	polyclonal	893
MF46.1	373	polyclonal	163	polyclonal	465	polyclonal	894
MF49.1	340	polyclonal	164	polyclonal	470	polyclonal	895
MF53.1	360	polyclonal	165	polyclonal	472	polyclonal	896
MF58.1	361	polyclonal	166	polyclonal	485	polyclonal	897
MF77.1	362	polyclonal	167	polyclonal	499	polyclonal	898
· · · · =		r J		r J		r J	

polyclonal	899	polyclonal	958	polyclonal	1277	T54
polyclonal	900	polyclonal	959	polyclonal	1280	T56
polyclonal	901	polyclonal	960	polyclonal	1281	T7.1
polyclonal	902	polyclonal	961	polyclonal	1282	T9
polyclonal	903	polyclonal	962	polyclonal	1285	T9
polyclonal	904	polyclonal	963	polyclonal	1296	TA9
polyclonal	905	polyclonal	964	polyclonal	1307	TB12
polyclonal	906	polyclonal	965	polyclonal	1349	TC15
polyclonal	907	polyclonal	966	polyclonal	1350	TD84
polyclonal	908	polyclonal	967	polyclonal	1351	TE135
polyclonal	909	polyclonal	968	polyclonal	1352	TG001
polyclonal	910	polyclonal	969	polyclonal	1353	TG002
polyclonal	911	polyclonal	970	polyclonal	1354	TH-Ab1
polyclonal	912	polyclonal	971	polyclonal	1355	TH1
polyclonal	913	polyclonal	972	polyclonal	1356	TH9
polyclonal	914	polyclonal	973	polyclonal	1357	V10
polyclonal	915	polyclonal	974	polyclonal	1358	V10-9
polyclonal	916	polyclonal	975	polyclonal	1359	V107
polyclonal	917	polyclonal	976	polyclonal	1360	V7-8
polyclonal	918	polyclonal	977	polyclonal	1361	W1
polyclonal	919	polyclonal	978	polyclonal	1362	W2
polyclonal	920	polyclonal	979	polyclonal	1363	X5
polyclonal	921	polyclonal	980	polyclonal	1364	Z13
polyclonal	922	polyclonal	981	polyclonal	1365	
polyclonal	923	polyclonal	982	polyclonal	1366	
polyclonal	924	polyclonal	983	polyclonal α577-596	647	
polyclonal	925	polyclonal	984	polyclonal α598-609	705	
polyclonal	926	polyclonal	985	polyclonal HIVIG	175	
polyclonal	927	polyclonal	986	RC25	541	
polyclonal	928	polyclonal	987	RL4.72.1	78	
polyclonal	929	polyclonal	988	RSD-33	394	
polyclonal	930	polyclonal	989	RT-4	249	
polyclonal	931	polyclonal	990	RT6H	196	
polyclonal	932	polyclonal	991	RT7O	250	
polyclonal	933	polyclonal	992	RT7U	251	
polyclonal	934	polyclonal	993	RTMAb8	194	
polyclonal	935	polyclonal	994	RV110026	626	
polyclonal	936	polyclonal	1000	S1-1	1106	
polyclonal	937	polyclonal	1020	SAR1	768	
polyclonal	938	polyclonal	1117	sc-FV p17	34	
polyclonal	939	polyclonal	1141	SC258	1155	
polyclonal	940	polyclonal	1199	SP.BAL114	500	
polyclonal	941	polyclonal	1200	SP.SF2:104	501	
polyclonal	942	polyclonal	1201	T1.1	341	
polyclonal	943	polyclonal	1202	T11	366	
polyclonal	944	polyclonal	1203	T13	1107	
polyclonal	945	polyclonal	1204	T15G1	869	
polyclonal	946	polyclonal	1205	T2.1	363	
polyclonal	947	polyclonal	1206	T20	870	
polyclonal	948	polyclonal	1207	T22	1128	
polyclonal	949	polyclonal	1212	T26	998	
polyclonal	950	polyclonal	1213	T27	871	
polyclonal	951	polyclonal	1214	T3	872	
polyclonal	952	polyclonal	1255	T30	873	
polyclonal	953	polyclonal	1266	T32	690	
polyclonal	954	polyclonal	1267	T34	691	
polyclonal	955	polyclonal	1271	T4	874	
polyclonal	956	polyclonal	1273	T49	1108	
polyclonal	957	polyclonal	1276	T52	1139	

# IV-B-3 MAbs by order of appearance in tables

No.	MAb ID	54	714/01	111	108/03	165	polyclonal
p17	WIAU ID	55	polyclonal	111	110/015	166	polyclonal
1	L14.17	56	111/073	113	32:32K	167	polyclonal
2	polyclonal	57	113/038	114	C5200	168	polyclonal
3	32/5.8.42	58	1-E-4	115	FH2	169	polyclonal
4	32/5.8.42	59	1-E-9	116	13B5	170	polyclonal
5	HyHIV-1	60	10-E-7	117	106/01	171	polyclonal
6	HyHIV-2	61	10-G-9	118	LH-104-B	172	polyclonal
7	HyHIV-3	62	11-C-5	119	LH-104-I	173	polyclonal
8	HyHIV-4	63	2-E-4	p24-j	p2p7p1p6	174	polyclonal
9	HyHIV-5	64	2-H-4	120	LH-104-G	175	polyclonal HIVIG
10	HyHIV-6	65	8-D-2	<b>p2p7</b>	p1p6	Prote	•
11	32/1.24.89	66	8-G-9	121	i5B11	176	1696
12	3B10	67	8-H-7	122	EC6	177	10E7
13	3E11	68	C5123	123	M12	178	F11.2.32
14	8H10	69	1-B-7	124	DG8	179	13E1
15	HyHIV-21	70	3-B-7	125	EB5	180	8B11
16	B4f8	71	6-D-12	126	НН3	181	8C10
17	HyHIV-22	72	6-E-7	127	AD2	182	8G5
18	12H-D3b3	73	8-D-5	128	CA5	RT	
19	12G-A8g2	74	FF1	129	DF3	183	1E8
20	12G-D7h11	75	113/072	130	EC3	184	polyclonal
21	12G-H1c7	76	25.3	131	FC12	185	1.152 B3
22	12I-D12g2	77	13-102-100	132	GE4	186	1.158 E2
23	polyclonal	78	RL4.72.1	133	JB7	187	31D6
24	HyHIV-15	79	406/01	134	JF11	188	31G8
25	11H9	80	polyclonal	Gag		189	32E7
26	3-H-7	81	38:9.6K	135	polyclonal	190	33D5
27	C5126	82	EB1A9	136	16/4/2	191	5B2
28	1D9	83	polyclonal	137	183-H12-5C	192	polyclonal
29	4C9	84	30:3E5	138	241-D	193	1.153 G10
30	4H2B1	85	EF7	139	2A6	194	RTMAb8
31	9G5	86	LH-104-E	140	5E2.A3k	195	1D4A3
32	15-21	87	1B2C12	141	71-31	196	RT6H
33	31-11	88	LH-104-K	142	91-6	197	1.160 B3
34	sc-FV p17	89	LH-104-A	143	98-4.3	198	polyclonal
թ17-լ		90	1.17.3	144	98-4.9	199	C2003
35	3A6	91	1A7	145	AC2	200	5B11
36	polyclonal	92	1F6	146	BC1071	201	6B10
p24		93	23A5G4	147	BE10	202	6E9
37	111/182	94	23A5G5	148	CD9	203	E-4
38	112/021	95	3D10G6	149	CH9B2	204	6B9
39	112/047	96	polyclonal	150	ED8	205	5F
40	ID8F6	97	F5-4	151	EH12E1	206	5G
41	F5-2	98	MO9.42.2	152	G11G1	207	7C4
42	CB-13/5	99	MO9.50.2	153	G11H3	Integ	
43	polyclonal	100	V10	154	HyHIV-19	208	1C4
44	3D3	101	V107	155	IE8G2	209	2C11
45	CD-4/1	102	LH-104-C	156	V7-8	210	2E3
46	15F8C7	103	12-B-4	157	anti-p24	211	3E11
47	111/052	104	C5122	158	human sera	212	3F9
48	polyclonal	105	9A4C4	159	polyclonal	213	5F8
49 50	91-5	106	11C10B10	160	polyclonal	214	6G5
50	1109/01	107	11D11F2	161	polyclonal	215	7B6
51	14D4E11	108	CD12B4	162	polyclonal	216	7C6
52 53	1G5C8	109	BE3	163	polyclonal	217	6C5
53	47-2	110	L14	164	polyclonal	218	8G4

219	17	274	polyclonal	330	L5.1	389	6C4/S
220	4D6	275	polyclonal	331	4A7C6	390	C108G
221	7-16	276	polyclonal	332	1D10	391	10/76b
222	4F6	277	1D2F11	333	B242	392	11/41e
223	anti-K159	278	2D9E7	334	133/192	393	11/4b
224	5D9	279	4B4C4	335	489.1(961)	394	RSD-33
225	8-6	280	5G7D8	336	5B3	395	11/4c
226	19	281	polyclonal	337	B10	396	8.22.2
227	2-19	282	NT2/4D5.24	338	B2	397	12b
228	8-22	283	polyclonal	339	C6	398	G3-136
229	4-20	284	I4: T-4	340	MF49.1	399	G3-4
230	6-19	285	L-anti-Tat	341	T1.1	400	BAT085
231 232	7C3 7F11	286 287	polyclonal	342 343	T7.1 T9	401 402	60b 74
			polyclonal				
233 234	8E5 MAb 35	288 289	polyclonal	344 345	GV4D3 B27	403 404	38/12b
235			polyclonal	343 346	B9	404	38/60b
Pol	polyclonal	290	polyclonal	340 347		406	polyclonal 322-151
	10	291	polyclonal		B35		
236	12	292	polyclonal	348	D/4B5 D/5A11	407	3D3.B8
237	13	293	2D9D5	349	D/5A11 D/6B2	408	4C11.D8
238	14	294	polyclonal	350		409	493-156
239	16 1C12B1	295	polyclonal	351	B18	410	110.1
240		296	polyclonal	352	B20	411	GV4H3
241	21	297	G1	353	MF39.1	412	J1
242	32	298	G2	354	187.2.1	413	J3
243 244	35 3D12	299 300	J1 TC15	355 356	37.1.1(ARP 327)	414 415	1006-30-D
					6D8		847-D
245	3F10 4	301 302	polyclonal	357 358	M96 MF119.1	416 417	MF169.1
246 247		303	polyclonal B1E3	359	MF4.1	417	MF170.1
247	6B9 7C4	303	J3B2	360	MF53.1	418	MF87.1 213.1
249	RT-4	Rev	J3D2	361	MF58.1	420	B12
250	RT7O	305	4G9	362	MF77.1	420	B12
251	RT7U	306	Ab2	363	T2.1	422	C13
252	anti-HIV-1 RT	307	10.1	364	11/65	422	M89
253		308	3H6	365	W1	424	B21
254	polyclonal	309	8E7	366	T11	424	B23
255	polyclonal	310	9G2	367	GV1A8	426	B23 B24
256	polyclonal 33	311	Ab4	368	11	427	B25
257	F-6	312	3G4	369	12G10	428	B23
Vif	1'-0	313	1G10	370	135/9	429	B26
258	TG002	314	1G7	371	7C10	430	B29
259	TG001	315	Ab3	372	C4	431	B36
260	J4	316	2G2	373	MF46.1	432	110.E
261	polyclonal	Vpu	202	374	6D5	433	110.E 110.C
Vpr	porycionar	317	DE7	375	B33	434	IIIB-V3-26
262	polyclonal	gp160		376	polyclonal	435	IIIB-V3-20 IIIB-V3-21
Tat	porycionar	318	M85	377	35D10/D2	436	168B8
263	polyclonal	319	7E2/4	378	40H2/C7	437	polyclonal
264	polyclonal	320	4D4#85	379	43A3/E4	438	polyclonal
265	TA9	321	M92	380	43C7/B9	439	MO97/V3
266	TD84	322	M86	381	45D1/B7	440	polyclonal
267	TE135	323	polyclonal	382	46E3/E6	441	55/11
268	polyclonal	324	133/237	383	58E1/B3	442	8/38c
269	NT3/2D1.1	325	133/290	384	64B9/A6	443	8/64b
270	1.2	326	133/11	385	69D2/A1	444	polyclonal
271	1D9D5	327	D/3G5	386	82D3/C3	445	polyclonal
272	TB12	328	D/6A11	387	2H1B	446	polyclonal
273	polyclonal	329	D/5E12	388	697-D	447	polyclonal
2,3	poljetonar	527	2,0212	200	V, D	,	Polycional

448	9284	507	G3-523	566	9205	625	43F
449	polyclonal	508	MN215	567	110.I	626	RV110026
450	polyclonal	509	Nea 9301	568	anti-HIV-2 polyclonal	627	105-306
451	polyclonal	510	4117C	569	IIIB-V3-01	628	GV1G2
452	polyclonal	511	419-D	570	D/6D1	629	750-D
453	MAG 109	512	453-D	571	4D7/4	630	450-D
454	MAG 49	513	504-D	572	36.1(ARP 329)	631	670-D
455	MAG 53	514	83.1	573	C12	632	158F3
456	MAG 56	515	5023B	574	110.D	633	161D7
457	1324-E	516	F58/D1	575	B32	634	polyclonal
458	polyclonal	517	P1/D12	576	polyclonal	635	722-D
459	MO99/V3	518	P4/D10	577	B15	636	polyclonal
460	C311E	519	IIIB-13 V3	578	B34	637	1331A
461	907	520	IIIB-34 V3	579	7F11	638	1131-A
462	924	521	A47/B1	580	E51	639	858-D
463	polyclonal	522	D59/A2	581	JL413	640	989-D
464	polyclonal	523	G44/H7	582	5C2E5	641	1A1
465	polyclonal	524	MO96/V3	583	G3-211	642	24G3
466	D19	525	μ5.5	584	G3-537	643	25C2
467	10F10	526	268-D	585	polyclonal	644	5F3
468	2C4	527	386-D	586	1795	645	$\alpha(566-586)$
469	412-D	528	5042A	587	ICR38.1a	646	PC5009
470	polyclonal	529	5042B	588	G3-299	647	polyclonal α577-596
471	CGP 47 439	530	418-D	589	G3-42	648	polyclonal
472	polyclonal	531	5021	590	G3-508	649	1.7
473	178.1	532	5025B	591	G3-519	650	
474	257-D	533	5042	592	G3-536	651	1F11
475	311-11-D	534	110.3	593	ICR38.8f	652	1H5
476	41148D	535	110.4	594	MO86/C3	653	3D9
477	391/95-D	536	110.5	595	13H8	654	4B3
478	Aw	537	58.2	596	G45-60	655	4D4
479	Bw	538	polyclonal	597	polyclonal	656	4G2
480	DO142-10	539	537-D	598	1662	657	polyclonal
481	Dv	540	5020	599	1663	658	polyclonal
482	Fv	541	RC25	600	1664	659	polytional
483	Gv	542	5023A	601	1697	660	polyclonal
484	Hv	543	110.6	602	1794	661	2A2/26
485	polyclonal	544	polyclonal	603	1804	662	50-69
486	50.1	545	10/36e	604	1807	663	9-11
487	30.1	546	10/54	605	1808	664	98-43
488	BAT123	547	11/85b	606	polyclonal	665	41-1
489	838-D	548	polyclonal	607	polyclonal	666	41.4
490	1006-15D	549	$0.5\beta$	608	CRA1	667	Fab A1
491	782-D	550	$C\beta 1, 0.5\beta$	609	M91	668	Fab A4
492	908-D	551	NM-01	610	9201	669	Fab M12B
493	1027-15D	552	1026	611	1C1	670	Fab M26B
494	F19.26-4	553	1034	612	3F5	671	Fab M8B
495	F19.48-3	554	59.1	613	5F4/1	672	Fab T2
496	F19.57-11	555	polyclonal	614	660-178	673	polyclonal
497	13105100	556	10E3	615	9301	674	86
498	M77	557	polyclonal	616	B221	675	polyclonal
499	polyclonal	558	N11-20	617	8C6/1	676	V10-9
500	SP.BAL114	559	5025A	618	H11	677	polyclonal
501	SP.SF2:104	560	N70-1.9b	619	W2	678	anti-P1
502	polyclonal	561	902	620	M38	679	polyclonal
503	19b	562	694/98-D	621	Chim 1	680	polyclonal
504	loop 2	563	MO101/V3,C4	622	polyclonal	681	2F11
505	4G10	564	MO101/V3,C4 MO101/V3,C4	623	110.1	682	246-D
506	5F7	565	MO101/V3,C4 MO101/V3,C4	624	42F	683	polyclonal
200	J1 1	505	1v1O101/ v J,C4	024	741	003	porycioliai

684	9G5A	743	41S-2	801	31710B	860	M25
685	181-D	744	447-52D	802	38B5/C9	861	MAG 6B
686	240-D	745	C8	803	39H10/A11	862	MO28
687	F240	746	B31	804	3C9	863	MO30
688	D49	747	B33	805	3D5	864	MO43
689	D61	748	1576	806	3H6	865	N2-4
690	T32	749	1578	807	40D3/C11	866	N70-2.3a
691	T34	750	1579	808	49B11/A1	867	P43110
692	115.8	751	1583	809	52G5/B9	868	P5-3
693	M-1	752	1899	810	55E4/H1	869	T15G1
694	M-11	753	1907	811	56C4/C8	870	T20
695	M-13	754	1908	812	57B6/F1	871	T27
696	M-2	755	1909	813	57H5/D7	872	T3
697	M-22	756	41-1	814	63G4/E2	873	T30
698	M-24	757	41-2	815	65B12/C5	874	T4
699	M-25	758	41-3	816	694/98D	875	m18
700	M-28	759	ED6	817	6D8	876	multiple Fabs
701	M-29	760	LA9 (121-134)	818	6E10	877	multiple MAbs
702	M-36	761	1575	819	7-1054	878	multiple MAbs
703	M-4	762	88-158/02	820	85G11/D8	879	multiple MAbs
704	M-6	763	88-158/022	821	87E4/A8	880	polyclonal
705	polyclonal α598-609	764	88-158/079	822	97B1/E8	881	polyclonal
706	1B8.env	765	polyclonal	823	A9	882	polyclonal
707	polyclonal	766	polyclonal	824	ADP421 polyclonal	883	polyclonal
708	polyclonal	767	B8	825	AG10H9	884	polyclonal
709	clone 3	768	SAR1	826	AH48	885	polyclonal
710	4	769	1577	827	B4	886	polyclonal
711	41-6	770	polyclonal	828	B5	887	polyclonal
712	41-7	771	DZ	829	B6	888	polyclonal
713	68.1	772	1010	830	BAT267	889	polyclonal
714	68.11	773	1014	831	BAT401	890	polyclonal
715	75	774	1018	832	BAT509	891	polyclonal
716	polyclonal	775	1019	833	C31	892	polyclonal
717	105-732	776	1020	834	CO11	893	polyclonal
718	3D6	777	1022	835	D1	894	polyclonal
719	F172-D8	778	1034	836	D12	895	polyclonal
720	D50	779	1492	837	D16	896	polyclonal
721	5-21-3	780	polyclonal	838	D4	897	polyclonal
722	120-16	781	polyclonal	839	D43	898	polyclonal
723	98-6	782	2F19C	840	F223	899	polyclonal
724	167-7	783	1334-D	841	F285	900	polyclonal
725	ND-15G1	Env		842	F424	901	polyclonal
726	167-D	784		843	F425 B4e8	902	polyclonal
727	polyclonal	785		844	F7	903	polyclonal
728	polyclonal	786		845	Fab A12	904	polyclonal
729	polyclonal	787		846	Fab A2	905	polyclonal
730	5B2	788		847	Fab L9	906	polyclonal
731	9G11	789	102-135	848	G12	907	polyclonal
732	TH-Ab1	790	1025	849	G2	908	polyclonal
733	polyclonal	791	105-134	850	H2	909	polyclonal
734	polyclonal	792	10E9	851	H8	910	polyclonal
735	polyclonal	793	126-50	852	HBW4	911	polyclonal
736	polyclonal	794	12H2	853	HIVIG	912	polyclonal
737	14D9	795	13.10	854	IVI-4G6	913	polyclonal
738	2F5	796	1B1	855	IgA6/30λ	914	polyclonal
739	4E10	797	1D10	856	IgA6/5k	915	polyclonal
740	Z13	798	1F7	857	IgA6/L4	916	polyclonal
741	B30	799	2G12	858	K14	917	polyclonal
742	polyclonal	800	30D	859	KU32	918	polyclonal
	- •						~ •

919	polyclonal	978	polyclonal	1037	21h	1096	M6
920	polyclonal	979	polyclonal	1038	28A11/B1	1097	MAG 116
921	polyclonal	980	polyclonal	1039	2G6	1098	MAG 12B
922	polyclonal	981	polyclonal	1040	35F3/E2		MAG 29B
923	polyclonal	982	polyclonal	1041	38G3/A9	1100	MAG 3B
924	polyclonal	983	polyclonal	1042	428	1101	
925	polyclonal	984	polyclonal	1043	448-D		MAG 72
926	polyclonal	985	polyclonal	1044	46D2/D5		MAG 86
927	polyclonal	986	polyclonal	1045	48-16	1104	MAG 96
928	polyclonal	987	polyclonal		50-61A		MTW61D
929	polyclonal	988	polyclonal	1047	5145A	1106	
930	polyclonal	989	polyclonal		558-D	1107	
931	polyclonal	990	polyclonal	1049	559/64-D	1108	T49
932	polyclonal	991	polyclonal	1050	55D5/F9	1109	T56
933	polyclonal	992	polyclonal	1051	588-D	1110	TH9
934	polyclonal	993	polyclonal	1052	654-D	1111	anti-CD4BS summary
935	polyclonal	994	polyclonal	1053	67G6/C4	1112	b11
936	polyclonal	995	101-342	1054	729-D	1113	b13
937	polyclonal	996	101-451	1055	830D	1114	b14
938	polyclonal	997	120-1	1056	9CL	1115	b3
939	polyclonal	998	T26	1057	BM12	1116	b6
940	polyclonal	999	D33	1058	D20	1117	polyclonal
941	polyclonal	1000	polyclonal	1059	D21	1118	
942	polyclonal	1001	212A	1060	D24	1119	17b
943	polyclonal	1002	522-149	1061	D25	1120	21c
944	polyclonal	1003	CA1	1062	D28	1121	23e
945	polyclonal	1004	CA13	1063	D35	1122	48d
946	polyclonal	1005	L19	1064	D39	1123	49e
947	polyclonal	1006	M90	1065	D42	1124	Fbb21
948	polyclonal	1007	MAG 104	1066	D52	1125	Fbb21
949	polyclonal	1008	MAG 45	1067	D53	1126	X5
950	polyclonal	1009	MAG 95	1068	D60	1127	8F101
951	polyclonal	1010	MAG 97	1069	DA48	1128	T22
952	polyclonal	1011	P35	1070	DO8i	1129	2A2
953	polyclonal	1012	T9	1071	F105	1130	AC4
954	polyclonal	1013	p7	1072	F91	1131	AD3
955	polyclonal		L100		FG39	1132	AD3
956	polyclonal		2/11c	1074	Fbb14	1133	
957	polyclonal	1016	A32	1075	GP13	1134	ID6
958	polyclonal	1017	C11	1076	GP44	1135	11/68b
959	polyclonal	1018	L81	1077	GP68	1136	62c
960	polyclonal	1019		1078	HF1.7	1137	CRA-6
961	polyclonal	1020	polyclonal	1079	HT5	1138	L15
962	polyclonal	1021	1024	1080	HT6	1139	T52
963	polyclonal	1022	4KG5	1081	HT7	1140	T54
964	polyclonal	1023	23A	1082	ICR 39.13g	1141	polyclonal
965	polyclonal	1024	D7324	1083	ICR 39.3b	1142	1088
966	polyclonal	1025	10/46c	1084	Ia3	1143	110-B
967	polyclonal	1026	1008-D	1085	Ia7	1144	1357
968	polyclonal	1027	1027-30-D	1086	IgG1b12	1145	1361
969	polyclonal	1028	1125H	1087	IgGCD4	1146	1393A
970	polyclonal	1029	1125H	1088		1147	2158
971	polyclonal		120-1B1	1089		1148	66a
972	polyclonal	1031	1202-D	1090	L41	1149	66c
973	polyclonal		1331E	1091			684-238
974	polyclonal		1570	1092			830A
975	polyclonal		1595	1093			CRA-3
976	polyclonal	1035	1599	1094	M12	1153	CRA-4
977	polyclonal	1036		1095	M13	1154	L17

1155	SC258	1214	polyclonal	1272	2E3	1330	2601
1156	L25	1215	2G12	1273	polyclonal	1331	2909
1157		1216	1367		F14.11	1332	4.11C
1158		1217	7B2	1275	31/03		4.6H
1159	L78	1218	126-6	1276	polyclonal	1334	4.8E
1160		1219	1342	1277	polyclonal		5145A
1161	10D8	1220	1379	1278	F4	1336	
1162	10F6	1221		1279			8.2A
1163	110.J		Fab D11		polyclonal		E047
	11G5		Fab D5	1281	polyclonal		ED10
	2182	1224	Fab G1		polyclonal		EH21
	2191	1225	Fab M10	1283	F3	1341	
1167	2219	1226	Fab M12	1284			F2A3
	2412		Fab M15	1285	polyclonal	1343	F3.9F
1169			Fab S10	1286			LA15
1170	2456	1229	Fab S6	1287	2F2	1345	LA21
1171	2483	1230	Fab S8	1288	E9	1346	LA28
1172	2497	1231	Fab S9	1289	3E6		LF17
	2557	1232	Fab T3	1290		1348	M2
	2558		Md-1	1291		1349	polyclonal
1175	2580		Fab A9	1292		1350	polyclonal
1176	391/95-D		Fab G15	1293		1351	polyclonal
1177	39F	1236	Fab G5	1294		1352	polyclonal
1178	4148d		Fab L1	1295	NF1A1	1353	polyclonal
1179	55/68b	1238	Fab L11		polyclonal	1354	polyclonal
1180	5G11		Fab L2	1297	E7	1355	polyclonal
1181	6.1	1240	1281	1298	AE6	1356	polyclonal
1182	6.7	1241	Chessie 8	1299	AG11	1357	polyclonal
1183	8.27.3	1242	8F102	1300	EH1	1358	polyclonal
1184	8E11/A8	1243	CG-10	1301	3B4B	1359	polyclonal
1185	9305		CG-25		3H3E	1360	polyclonal
1186	A1g8		CG-4	1303		1361	polyclonal
1187	AG1121		CG-76		NF2B2	1362	polyclonal
1188	Ag1211		CG-9		NF3A3	1363	polyclonal
1189		1248	105-518		NF8B4	1364	polyclonal
1190	B4e8	1249	31A1		polyclonal	1365	polyclonal
1191	D27	1250	39A64	1308		1366	polyclonal
1192		1251	39B86	HIV-	1		
1193		1252	9303	1309			
1194			NC-1	1310			
	G3-1472	Nef		1311			
1196		1254			1.4C		
1197			polyclonal		1.4G		
1198	anti-gp120/V3		13/042		1.9E		
1199	polyclonal		13/035		1.9F		
1200	polyclonal	1258			12.19		
1201	polyclonal		AM5C6		12.9		
1202	1 2		AM5C6		13a15		
1203	polyclonal	1261			13a23		
1204	polyclonal		25/03		13a3		
1205	polyclonal		26/76		13a6		
1206	1 3	1264			13a7		
1207	polyclonal		3D12		13b18		
1208	11/75a/21/41		polyclonal		13b23		
1209			polyclonal	1325	13b53		
	55/45a/11		3G12		13b61		
1211	1108		13/058		2.5E		
	polyclonal	1270			2191		
1213	polyclonal	1271	polyclonal	1329	25G		

# IV-C

# **HIV Antibodies Tables**

All HIV MAbs and polyclonal Abs that bind to linear epitopes 30 amino acids or less in length are arranged by protein position. The table entries are sorted in a nested way—first by protein, then by HXB2 start location, then by antibody type, and finally by antibody name. Abs that bind to conformational epitopes or with unknown epitopes are listed at the end of each protein section.

## IV-C-1 Gag p17 Antibodies

No. 1

**MAb ID** L14.17

**HXB2 Location** p17 (11-25)

Author Location p17 (11-25 BRU)

Epitope GELDRWEKIRLRPGG

Neutralizing no

Immunogen vaccine

Vector/Type: viral lysate Strain: B clade

BRU HIV component: HIV-1

Species (Isotype) mouse (IgG)

References Robert-Hebmann et al. 1992a; Robert-

Hebmann et al. 1992b; Tatsumi et al. 1990

**No.** 2

MAb ID polyclonal

HXB2 Location p17 (11-25)

**Author Location** p17 (11–25 LAI)

Epitope GELDRWEKIRLRPGG

Subtype B

Neutralizing no

Immunogen vaccine

*Vector/Type:* protein, virus-like particle (VLP) Strain: B clade LAI HIV component: Gag, p17 Gag, p24 Gag Adjuvant: Complete

Freund's Adjuvant (CFA)

Species (Isotype) mouse

References Truong et al. 1997

• An ELISA assay was used to study a panel of Gag peptides mature p24 CA epitopes mapped to residues 176-192, 201-218, 233-253, 285-304, and were recognized by antibodies elicited by rp24CA – one p17MA epitope, residues 11-25, and one p24CA epitope, residues 176-192, were recognized by antibodies raised against anti-p55 virus-like particles, suggesting a differentantigenic properties for p24CA and p17MA antibodies depending on whether they are produced against the mature soluble protein or the immature assembled form of the gag proteins. Truong et al. [1997]

> **No.** 3 MAb ID 32/5.8.42

**HXB2 Location** p17 (12–19)

Author Location p17 (12–19 IIIB)

Epitope ELDRWEKI+ALDKIE

Neutralizing no

Immunogen vaccine

Vector/Type: viral lysate

Species (Isotype) mouse (IgG)

References Papsidero et al. 1989

• 32/5.8.42: Binds to two discontinuous regions, positions 12-19 and 100-105, peptides ELDRWEKI and ALDKIE - inhibited infectivity of cell free virus. Papsidero et al. [1989]

No. 4

**MAb ID** 32/5.8.42

**HXB2 Location** p17 (12–19)

Author Location p17 (IIIB)

Epitope ELDRWEKI+ALDKIE

Neutralizing no

Immunogen vaccine

Vector/Type: viral lysate HIV component:

Species (Isotype) mouse (IgG)

References Papsidero et al. 1989

• 32/5.8.42: Inhibited infectivity of cell free virus – bound to two peptides, ELDRWEKI and ALDKIE, at positions 12-19 + 100-105. Papsidero et al. [1989]

**No.** 5

MAb ID HyHIV-1

**HXB2 Location** p17 (12–29)

Author Location p17 (12–29 JMH1)

Epitope ELDKWEKIRLRPGGKTLY

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

Species (Isotype) mouse (IgG1)

References Ota & Ueda 1998; Liu et al. 1995

• HyHIV-1: This paper compares the results of affinity constant (Ka) measurements of anti-p17 MAbs using double Ab methods versus the faster, isotope-free BIAcore system, and results were found to be similar for HyHIV-(1-6) – six MAbs all bind to the first alpha helix of p17, a functional domain for both membrane binding and nuclear localization. Ota & Ueda [1998]

**No.** 6

MAb ID HyHIV-2

HXB2 Location p17 (12-29)

Author Location p17 (12–29 JMH1)

Epitope ELDKWEKIRLRPGGKTLY

HIV Antibodies Tables Gag p17 Antibodies

Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

Species (Isotype) mouse (IgG1)

References Ota & Ueda 1998; Liu et al. 1995

• HyHIV-2: This paper compares the results of affinity constant (Ka) measurements of anti-p17 MAbs using double Ab methods versus the faster, isotope-free BIAcore system, and results were found to be similar for HyHIV-(1-6) – six MAbs all bind to the first alpha helix of p17, a functional domain for both membrane binding and nuclear localization. Ota & Ueda [1998]

**No.** 7

MAb ID HyHIV-3

**HXB2 Location** p17 (12–29)

Author Location p17 (12-29 JMH1)

Epitope ELDKWEKIRLRPGGKTLY

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

Species (Isotype) mouse (IgG1)

References Ota & Ueda 1998; Liu et al. 1995

• HyHIV-3: This paper compares the results of affinity constant (Ka) measurements of anti-p17 MAbs using double Ab methods versus the faster, isotope-free BIAcore system, and results were found to be similar for HyHIV-(1-6) – six MAbs all bind to the first alpha helix of p17, a functional domain for both membrane binding and nuclear localization. Ota & Ueda [1998]

**No.** 8

MAb ID HyHIV-4

**HXB2 Location** p17 (12–29)

Author Location p17 (12–29 JMH1)

Epitope ELDKWEKIRLRPGGKTLY?

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

**Species (Isotype)** mouse (IgG1)

**References** Ota & Ueda 1998; Ota *et al.* 1998; Liu *et al.* 1995

- HyHIV-4: epitope uncertain, based on the best estimate from JMH1 sequence— Ka is 1.8 x 10<sup>7</sup> M-1 for rec p17 stains the surface of infected cells indicating the antigen is exposed at the cell surface. Ota *et al.* [1998]
- HyHIV-4: This paper compares the results of affinity constant (Ka) measurements of anti-p17 MAbs using double Ab methods versus the faster, isotope-free BIAcore system, and results were found to be similar for HyHIV-(1-6) six MAbs all bind to the first alpha helix of p17, a functional domain for both membrane binding and nuclear localization. Ota & Ueda [1998]

No. 9

MAb ID HyHIV-5

HXB2 Location p17 (12-29)

**Author Location** p17 (12–29 JMH1)

Epitope ELDKWEKIRLRPGGKTLY

Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

**Species (Isotype)** mouse (IgG1)

References Ota & Ueda 1998; Liu et al. 1995

HyHIV-5: This paper compares the results of affinity constant (Ka) measurements of anti-p17 MAbs using double Ab methods versus the faster, isotope-free BIAcore system, and results were found to be similar for HyHIV-(1-6) – six MAbs all bind to the first alpha helix of p17, a functional domain for both membrane binding and nuclear localization. Ota & Ueda [1998]

**No.** 10

MAb ID HyHIV-6

HXB2 Location p17 (12-29)

**Author Location** p17 (12–29 JMH1)

Epitope ELDKWEKIRLRPGGKTLY

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

Species (Isotype) mouse (IgG1)

References Ota & Ueda 1998; Liu et al. 1995

• HyHIV-6: This paper compares the results of affinity constant (Ka) measurements of anti-p17 MAbs using double Ab methods versus the faster, isotope-free BIAcore system, and results were found to be similar for HyHIV-(1-6) – six MAbs all bind to the first alpha helix of p17, a functional domain for both membrane binding and nuclear localization. Ota & Ueda [1998]

No. 11

MAb ID 32/1.24.89

HXB2 Location p17 (17-22)

**Author Location** p17 (17–22 IIIB)

Epitope EKIRLR

Neutralizing L

Immunogen vaccine

Vector/Type: viral lysate

Species (Isotype) mouse (IgG)

References Papsidero et al. 1989

• 32/1.24.89: Inhibited infectivity of cell free virus. Papsidero *et al.* [1989]

**No.** 12

**MAb ID** 3B10

**HXB2 Location** p17 (19–38)

**Author Location** p17 (19–38 SIVmac)

Epitope IRLPGGKKKYMLKHVVWAA

Neutralizing no

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade AGM TYO-7 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Otteken et al. 1992

Gag p17 Antibodies **HIV Antibodies Tables** 

• 3B10: Recognized an epitope present on HIV-2/SIVmac (MAC251/32H), SIVagm, HIV-1, and SIVmnd, demonstrating that the matrix protein of all nine HIV and SIV isolates tested in this study expresses at least one conserved immunogenic epitope recognized serologically. Otteken et al. [1992]

**No.** 13 MAb ID 3E11 HXB2 Location p17 (19-38) **Author Location** p17 (19–38 SIVmac) Epitope IRLPGGKKKYMLKHVVWAA

> Neutralizing no Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade AGM TYO-7 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Nilsen et al. 1996; Otteken et al. 1992

- 3E11: There is another MAb with this ID that recognizes integrase. Nilsen et al. [1996]
- 3E11: Recognized an epitope present on HIV-2/SIVmac (MAC251/32H), SIVagm, HIV-1, and SIVmnd, demonstrating that the matrix protein of all nine HIV and SIV isolates tested in this study expresses at least one highly conserved immunogenic epitope. Otteken et al. [1992]

No. 14 MAb ID 8H10 **HXB2 Location** p17 (30–52) Author Location p17 (30-52 JMH1)

Epitope KLKHIVWASRELERFAVNPGLLE

Neutralizing

Immunogen vaccine

*Vector/Type:* peptide *Strain:* B clade JMH-1 HIV component: p17 Gag Adjuvant: BSA

Species (Isotype) mouse (IgM)

References Ota & Ueda 1999; Ota et al. 1999

- 8H10: The p17 MAb also can bind to the V3 loop. Ota et al.
- 8H10: Inhibits viral replication of the HIV-1 infected MT-4 cells by decreasing p17 DNA levels in the infected cells, and the effect of growing the 8H10 hybridoma in co-culture with HIV-1 infected MT-4 cells was studied. Ota & Ueda [1999]

No. 15

MAb ID HyHIV-21

HXB2 Location p17 (30-52)

Author Location p17 (30–52 JMH1)

Epitope KLKHIIWASRELERFAVNPGLLE

Neutralizing no Immunogen vaccine

> Vector/Type: protein HIV component: p17 Gag

Species (Isotype) mouse (IgG2a)

References Ota et al. 1998; Liu et al. 1995

• HyHIV-21: epitope uncertain, based on the best estimate from JMH1 sequence – Ka is  $3.6 \times 10^6$  M-1 for rec p17 – stains the surface of infected cells indicating the antigen is exposed at the cell surface -inhibited growth of HIV-1 JMH1 in MT-4 cells when added 24 hours after the initial culture. Ota et al. [1998]

**No.** 16

MAb ID B4f8

**HXB2 Location** p17 (51–65)

Author Location p17 (51-65)

Epitope LETSEGCRQILGQLQ

Neutralizing no

Immunogen vaccine

HIV infected-cell lysate *Vector/Type:* Strain: B clade IIIB HIV component:

HIV-1

Species (Isotype) rat (IgG2a)

References Shang et al. 1991

• -B4f8: Did not bind live infected cells, only cells that had been made permeable with acetone. Shang et al. [1991]

No. 17

MAb ID HyHIV-22

HXB2 Location p17 (52-83)

**Author Location** p17 (53–87 JMH1)

Epitope ETSEGCRQILGQRQPSLQTGSEELRSLYNTIH

Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

Species (Isotype) mouse (IgG1)

References Ota et al. 1998; Liu et al. 1995

• HyHIV-22: epitope uncertain, based on the best estimate from JMH1 sequence – stains the surface of infected cells indicating the antigen is exposed at the cell surface – Ka is 2.3 x 10<sup>5</sup> M-1 for rec p17. Ota et al. [1998]

**No.** 18

**MAb ID** 12H-D3b3

**HXB2 Location** p17 (62–78)

Author Location p17 (62–78)

Epitope GQLQPSLQTGSEELRSL

Neutralizing no

Immunogen vaccine

Vector/Type: HIV infected-cell lysate Strain: B clade IIIB HIV component:

HIV-1

Species (Isotype) rat (IgG2a)

References Shang et al. 1991

• 12H-D3b3: Did not bind live infected cells, only cells that had been made permeable with acetone. Shang et al. [1991]

**No.** 19

MAb ID 12G-A8g2

HXB2 Location p17 (86-115)

**Author Location** p17 (86–115)

Epitope YCVHQRIEIKDTKEALDKIEEEQNKSKKKA

Neutralizing no Immunogen vaccine

> HIV infected-cell lysate Vector/Type: Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) rat (IgG2a)

References Maksiutov et al. 2002; Shang et al. 1991

HIV Antibodies Tables Gag p17 Antibodies

 12G-A8g2: This epitope is similar to a fragment of the human protein CD40 ligand TNF-related activation protein(T-cell antigen GP39) (CD154), LDKIEDERN, as well as to a fragment of Lens-epithelium-derived growth factor, DIITEEDKSKKKGQ. Maksiutov et al. [2002]

 12G-A8g2: Bound to 30-mer, but not to internal peptides – did not bind live infected cells – antigenic domain known as HPG30. Shang *et al.* [1991]

No. 20

MAb ID 12G-D7h11

HXB2 Location p17 (86-115)

Author Location p17 (86-115)

Epitope YCVHQRIEIKDTKEALDKIEEEQNKSKKKA

Neutralizing no

Immunogen vaccine

Vector/Type: HIV infected-cell lysate Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) rat (IgG2a)

References Maksiutov et al. 2002; Shang et al. 1991

- 12G-D7h11: This epitope is similar to a fragment of the human protein CD40 ligand TNF-related activation protein (T-cell antigen GP39) (CD154), LDKIEDERN, as well as to a fragment of Lens-epithelium-derived growth factor, DIITEEDKSKKKGQ. Maksiutov et al. [2002]
- 12G-D7h11: Bound to 30-mer, but not to internal peptides did not bind live infected cells antigenic domain known as HPG30. Shang *et al.* [1991]

**No.** 21

**MAb ID** 12G-H1c7

**HXB2 Location** p17 (86–115)

Author Location p17 (86–115)

Epitope YCVHQRIEIKDTKEALDKIEEEQNKSKKKA

Neutralizing no

Immunogen vaccine

Vector/Type: HIV infected-cell lysate Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) rat (IgG)

References Maksiutov et al. 2002; Shang et al. 1991

- 12G-H1c7: This epitope is similar to a fragment of the human protein CD40 ligand TNF-related activation protein (T-cell antigen GP39) (CD154), LDKIEDERN, as well as to a fragment of Lens-epithelium-derived growth factor, DIITEEDKSKKKGQ. Maksiutov *et al.* [2002]
- 12G-H1c7: Bound to 30-mer, but not to internal peptides did not bind live infected cells – antigenic domain known as HPG30. Shang et al. [1991]

No. 22

MAb ID 12I-D12g2

HXB2 Location p17 (86-115)

**Author Location** p17 (86–115)

Epitope YCVHQRIEIKDTKEALDKIEEEQNKSKKKA

Neutralizing no

Immunogen vaccine

Vector/Type: HIV infected-cell lysate Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) rat (IgG2a)

References Maksiutov et al. 2002; Shang et al. 1991

- 12I-D12g2: This epitope is similar to a fragment of the human protein CD40 ligand TNF-related activation protein (T-cell antigen GP39) (CD154), LDKIEDERN, as well as to a fragment of Lens-epithelium-derived growth factor, DIITEEDKSKKKGQ. Maksiutov *et al.* [2002]
- 12I-D12g2: Bound to 30-mer, but not to internal peptides did not bind live infected cells antigenic domain known as HPG30. Shang *et al.* [1991]

**No.** 23

MAb ID polyclonal

HXB2 Location p17 (86-115)

**Author Location** p17 (86–115)

Epitope YSVHQRIDVKDTKEALEKIEEEQNKSKKKA

Neutralizing L

Immunogen vaccine

Vector/Type: peptide HIV component: p17

Gag Adjuvant: Cholera toxin (CT)

**Species (Isotype)** mouse (IgA) **References** Bukawa *et al.* 1995

Polyclonal secretory IgA antibody raised by oral mucosal immunization is able to neutralize IIIB, SF2, and MN – HIV-1 neutralization may be due to the V3, CD4 or HPG30 component of the multicomponent peptide immunogen. Bukawa *et al.* [1995]

No. 24

MAb ID HyHIV-15

**HXB2 Location** p17 (87–115)

**Author Location** p17 (87–115 JMH1)

Epitope SVHQRIDVKDTKEALEKIEEEQNKSKKKA?

Neutralizing L

Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

Species (Isotype) mouse (IgG1)

References Ota et al. 1998; Liu et al. 1995

 HyHIV-15: epitope uncertain, based on the best estimate from JMH1 sequence – Ka is 1.4 x 10<sup>7</sup> M-1 for rec p17 – stains the surface of infected cells indicating the antigen is exposed at the cell surface – inhibited growth of HIV-1 JMH1 in MT-4 cells when added 24 hours after the initial culture. Ota et al. [1998]

No. 25

**MAb ID** 11H9

**HXB2 Location** p17 (101–115)

Author Location p17 (101–115 SF2)

**Epitope** LEKIEEEQNKSKKKA?

Neutralizing

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B

clade CBL-1 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

Research Contact R. B. Ferns and R. S. Tedder

Gag p17 Antibodies **HIV Antibodies Tables** 

References Maksiutov et al. 2002; Ferns et al. 1989; Ferns et al. 1987

- 11H9: UK Medical Research Council AIDS reagent: ARP344.
- 11H9: This epitope is similar to a fragment of Lens-epitheliumderived growth factor, DIITEEDKSKKKGQ. Maksiutov et al. [2002]
- 11H9: Reactive against p18 and p55. Ferns et al. [1987]

No. 26

**MAb ID** 3-H-7 (3H7)

HXB2 Location p17 (113-122)

Author Location p17 (113-122 BH10)

Epitope KKAQQAAADT

Neutralizing L

Immunogen vaccine

Strain: B clade IIIB

Species (Isotype) mouse (IgG)

References Levin et al. 1997; Robert-Hebmann et al. 1992a; Robert-Hebmann et al. 1992b; Niedrig et al. 1989

- 3-H-7: Called 3H7 using a bicistronic vector, an intracellular Fab intrabody, 3H7, can inhibit HIV-1 infection when expressed in the cytoplasm of dividing CD4+ T cells - HXBI-IIB and SI primary isolate virions from 3H7 expressing cells Research Contact R. B. Ferns and R. S. Tedder were far less infectious – 3H7 intrabody acts both at the stage of nuclear import and virus particle assembly. Levin et al. [1997]
- 3-H-7: No cross-reactivity with HIV-2 ROD or SIV MAC by immunoblot. Niedrig et al. [1989]

No. 27

MAb ID C5126

HXB2 Location p17 (113-122)

Author Location p17 (113–122 HXB2)

Epitope KKAQQAAADT

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: viral lysate HIV component:

HIV-1

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Hinkula et al. 1990

• C5126: Defined epitope by peptide blocking of binding to native protein – WB reactive with p53 and p17. Hinkula et al. [1990]

No. 28

MAb ID 1D9

HXB2 Location p17 (119-132)

Author Location p17 (121–134 SF2)

Epitope AAGTGNSSQVSQNY

**Neutralizing** 

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade CBL-1 HIV component: HIV-1

Species (Isotype) mouse (IgG2a)

Research Contact R. B. Ferns and R. S. Tedder

References Ferns et al. 1989; Ferns et al. 1987

- 1D9: UK Medical Research Council AIDS reagent: ARP316.
- 1D9: Reactive against p18, but not p55. Ferns et al. [1987]

No. 29

MAb ID 4C9

**HXB2 Location** p17 (119–132)

Author Location p18 (121-134 SF2)

**Epitope** AAGTGNSSQVSQNY

Neutralizing

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B

clade CBL-1 HIV component: HIV-1

Species (Isotype) mouse (IgG2a)

Research Contact R. B. Ferns and R. S. Tedder

References Ferns et al. 1989; Ferns et al. 1987

- 4C9: UK Medical Research Council AIDS reagent: ARP342.
- 4C9: Reactive against p18, but not p55. Ferns et al. [1987]

**No.** 30

MAb ID 4H2B1

HXB2 Location p17 (119-132)

Author Location p17 (121–134 SF2)

**Epitope** AAGTGNSSQVSQNY

**Neutralizing** 

Immunogen

Species (Isotype) mouse (IgG1)

References Ferns et al. 1989; Ferns et al. 1987

- 4H2B1: UK Medical Research Council AIDS reagent: ARP315.
- 4H2B1: Reactive against p18 and p55 of multiple isolates. Ferns et al. [1987]

No. 31

MAb ID 9G5

**HXB2 Location** p17 (119–132)

Author Location p17 (121–134 SF2)

Epitope AAGTGNSSQVSQNY

Neutralizing

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade CBL-1 HIV component: HIV-1

Species (Isotype) mouse (IgM)

Research Contact R. B. Ferns and R. S. Tedder

References Ferns et al. 1989; Ferns et al. 1987

- 9G5: UK Medical Research Council AIDS reagent: ARP343.
- 9G5: Reactive against p18, but not p55. Ferns et al. [1987]

**No.** 32

MAb ID 15-21

**HXB2 Location** p17 (121–132)

Author Location p17 (121–132 BRU)

**Epitope** DTGHSSQVSQNY

Neutralizing no

Immunogen vaccine

Strain: B clade BRU

**Species (Isotype)** mouse (IgG)

References Robert-Hebmann et al. 1992a; Robert-

Hebmann et al. 1992b

**No.** 33

**MAb ID** 31-11

**HXB2 Location** p17 (121–132)

**Author Location** p17 (121–132 BRU) **Epitope** DTGHSSQVSQNY

Neutralizing no Immunogen vaccine

Strain: B clade BRU

Species (Isotype) mouse (IgG)

**References** Robert-Hebmann *et al.* 1992a; Robert-Hebmann *et al.* 1992b

**No.** 34

**MAb ID** sc-FV p17 **HXB2 Location** p17 (121–132)

Author Location p17 (121-132 BRU)

Epitope DTGHSSQVSQNY

**Neutralizing** L

Immunogen vaccine

Strain: B clade BRU

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type C-term

Research Contact Paul Zhou, NIH, Bethesda, MD, USA

**References** Tewari *et al.* 1998; Robert-Hebmann *et al.* 1992a

 A single chain Ab (sc-FV) was made from an anti-p17 MAb, and intracellular binding of sc-FV resulted in inhibition of viral replication that was more pronounced when the sc-FV was expressed in the cytoplasm instead of the nucleus. Tewari *et al*. [1998]

## IV-C-2 Gag p17-p24 Antibodies

**No.** 35

MAb ID 3A6

**HXB2 Location** p17-p24 (122–17) **Author Location** p24 (122–149 BH10)

Epitope TGHSSQVSQNYPIVQNIQGQMVHQAISP

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

References Buchacher et al. 1994; Buchacher et al. 1992

- 3A6: The reactive peptide spans the p17/p24 border of gag. Buchacher *et al.* [1994]
- 3A6: Human MAbs against HIV generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994]

**No.** 36

MAb ID polyclonal

HXB2 Location p17-p24

**Author Location** Gag

Epitope

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Lottersberger et al. 2004

Addition of non-immunogenic side chains (AAAC and CAAA) to both N- and C-termini of synthetic alpha helical peptide sequences of HIV-1 p24 and p17 proteins improved Ab reactivity. As diminishing response to these antigens is a harbinger of progression these peptides may by useful in diagnostic assays. Lottersberger et al. [2004]

#### IV-C-3 Gag p24 Antibodies

**No.** 37

MAb ID 111/182

**HXB2 Location** p24 (1-20)

Author Location p24 (134–153 IIIB)

Epitope PIVQNIQGQMVHQAISPRTL

Neutralizing no

Immunogen vaccine

Vector/Type: beta-galactosidase fusion protein Strain: B clade IIIB HIV component:

p24 Gag

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1991

• 111/182: Test specific evidence of cross-reactivity between HIV-1, HIV-2 and SIV MAC. Niedrig *et al.* [1991]

**No.** 38

MAb ID 112/021

**HXB2 Location** p24 (1–20)

**Author Location** p24 (134–153 IIIB)

Epitope PIVQNIQGQMVHQAISPRTL

Neutralizing no

Immunogen vaccine

*Vector/Type:* beta-galactosidase fusion protein *Strain:* B clade IIIB *HIV component:* 

p24 Gag

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1991

• 112/021: Test specific evidence of cross-reactivity between HIV-1, HIV-2 and SIV MAC. Niedrig *et al.* [1991]

**No.** 39

MAb ID 112/047

**HXB2 Location** p24 (1-20)

Author Location p24 (134–153 IIIB)

Epitope PIVQNIQGQMVHQAISPRTL

Neutralizing no

Immunogen vaccine

Vector/Type: beta-galactosidase fusion protein Strain: B clade IIIB HIV component:

p24 Gag

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1991

• 112/047: Test specific evidence of cross-reactivity between HIV-1, HIV-2 and SIV MAC. Niedrig *et al.* [1991]

**No.** 40

MAb ID ID8F6

**HXB2 Location** p24 (11–25)

**Author Location** p24 (143–157 BRU)

Epitope VHQAISPRTLNAWVK

Gag p24 Antibodies **HIV Antibodies Tables** 

Neutralizing no Immunogen vaccine

> Vector/Type: inactivated HIV Strain: B clade CBL-1 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

Research Contact R. B. Ferns and R. S. Tedder

References Ferns et al. 1989; Ferns et al. 1987

- ID8F6: UK Medical Research Council AIDS reagent:
- ID8F6: Reacted with both p55 and p24 showed less than 75% homologous inhibition. Ferns et al. [1987]

**No.** 41

MAb ID F5-2

HXB2 Location p24 (14-23)

**Author Location** p24 (14–23 HXB2)

**Epitope** AISPRTLNAW

Subtype B

Neutralizing no

Immunogen

Species (Isotype) mouse

References Kusk et al. 1992; Kusk et al. 1988

• F5-2: In HIV-1 + individuals, antibody to AISPRTLNAW is Species (Isotype) mouse (IgG2b) associated with CD4 T-cell decline. Kusk et al. [1988, 1992] Research Contact R. B. Ferns and R. S. Tedder

No. 42

**MAb ID** CB-13/5 (CB-mab-p24/13-15)

**HXB2 Location** p24 (21–25)

**Author Location** p24 (152–156)

**Epitope** NAWVK

Neutralizing no

Immunogen

**Species (Isotype)** mouse (IgG1 $\kappa$ )

**References** Glaser & Hausdorf 1996: Kuttner et al. 1992: Franke et al. 1992; Grunow et al. 1990

- CB-13/5: It is not clear whether the MAbs CD-13/5 and CBmab-p24/13-15 are the same, but from the shared references in the primary articles they seem to be (database note)
- CB-13/5: Epitope described as VHQAISPRTLNAWVK binding not affected by bound MAb CB-4/1. Glaser & Hausdorf [1996]
- CB-13/5: Inhibits spread of HIV-1 in cell cultures. Franke et al.
- CB-13/5: Called CB-mab-p24/13-15 the VDJ H and VJ L regions of CB-mab-p24/13-15 were sequenced. Kuttner et al. [1992]

**No.** 43

MAb ID polyclonal

HXB2 Location p24 (44-60)

**Author Location** p24 (176–192 LAI)

Epitope SEGATPQDLNTMLNTVG

Subtype B

Neutralizing no

Immunogen vaccine

*Vector/Type:* protein, virus-like particle (VLP) Strain: B clade LAI HIV component: Gag, p17 Gag, p24 Gag Adjuvant: Complete Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgG)

References Truong et al. 1997

• An ELISA assay was used to study a panel of Gag peptides – mature p24 CA epitopes mapped to residues 176-192, 201-218, 233-253, 285-304, and were recognized by antibodies elicited by rp24CA - one p17MA epitope, residues 11-25, and one p24CA epitope, residues 176-192, were recognized by antibodies raised against anti-p55 virus-like particles, suggesting a different antigenic properties for p24CA and p17MA antibodies depending on whether they are produced against the mature soluble protein or the immature assembled form of the gag proteins. Truong et al. [1997]

No. 44

MAb ID 3D3

**HXB2 Location** p24 (45–50)

Author Location p24 (177-182 LAI)

**Epitope** EGATPQ

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade CBL-1 HIV component: HIV-1

References Ferns et al. 1989; Ferns et al. 1987

- 3D3: UK Medical Research Council AIDS reagent: ARP314.
- 3D3: Most broadly reactive of all the antibodies in this study. Ferns et al. [1987]

No. 45

MAb ID CD-4/1 (CB-4/1/1/F6)

**HXB2** Location p24 (46–56)

**Author Location** p24 (182–197)

Epitope GATPQDLNTML

Neutralizing no

Immunogen vaccine

Vector/Type: beta-galactosidase fusion pro-

tein HIV component: p24 Gag

Species (Isotype) mouse (IgG2a $\kappa$ )

References Ehrhard et al. 1996: Glaser & Hausdorf 1996: Hohne et al. 1993; Franke et al. 1992; Grunow et al. 1990

- CD-4/1: Modification of p24 lysine residues by maleic anhydrid increased the affinity of CD-4/1, presumably due to conformational changes exposing a cryptic epitope. Ehrhard et al. [1996]
- CD-4/1: Unusual p24-MAb binding kinetics, with biphasic association – probably due to conformational changes in p24, not to p24 dimerization. Glaser & Hausdorf [1996]
- CD-4/1: Affinity of CB-4/1 to native p24 is lower than to peptide or denatured p24 – proposed that the peptide binds in a loop conformation. Hohne et al. [1993]
- CD-4/1: Inhibits spread of HIV-1 in cell cultures. Franke et al. [1992]

No. 46

MAb ID 15F8C7

**HXB2 Location** p24 (47–56)

**Author Location** p24 (183–197)

Epitope ATPQDLNTML

HIV Antibodies Tables Gag p24 Antibodies

Neutralizing no

Immunogen vaccine

Vector/Type: purified HIV-1

Species (Isotype) mouse (IgG1)

References Janvier et al. 1992; Janvier et al. 1990

• 15F8C7: Mapped to aa209-217 through Pepscan method – cross-reacts with HIV-2 Janvier *et al.* [1990] – maps to aa203-217 through EIA pentadecapeptide Janvier *et al.* [1992]. Janvier *et al.* [1990, 1992]

No. 47

MAb ID 111/052

HXB2 Location p24 (51-60)

Author Location p24 (183–192 IIIB)

Epitope DLNTMLNTVG

Neutralizing no

Immunogen vaccine

Vector/Type: beta-galactosidase fusion protein Strain: B clade IIIB HIV component:

p24 Gag

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1991

• 111/052: Weak cross-reaction with HIV-2 on WB, otherwise not cross-reactive with HIV-2 or SIV MAC. Niedrig *et al.* [1991]

No. 48

MAb ID polyclonal

**HXB2 Location** p24 (51–82)

Author Location Gag (183-214 LAI)

Epitope DLNTMLNTVGGHQAAMQMLKETINEEAAEWDR

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: p24 Gag Adjuvant:

QS21

Species (Isotype) human (IgG)

References Pialoux et al. 2001

• 28 subjects were vaccinated with six HIV-1 peptides that were selected to be particularly rich in CTL epitopes, presented in lipopeptides with or without adjuvant QS21 – HIV-specific Ab responses were detected in 25/28 (89%), proliferative in 19/28 (79%), and CTL in 13/24 (54%) of testable volunteers – only 4/28 had Ab responses to peptide G1, 4/28 had proliferative responses, and no patient had a CTL response. Pialoux *et al.* [2001]

**No.** 49

**MAb ID** 91-5

**HXB2 Location** p24 (64–75)

**Author Location** p24 (196–207)

Epitope AAMQMLKETINE

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

References Gorny et al. 1998; Robinson et al. 1990b;

Tyler et al. 1990; Gorny et al. 1989

 91-5: NIH AIDS Research and Reference Reagent Program: 1238. • 91-5: Did not enhance HIV-1 IIIB infection. Robinson *et al.* [1990b]

 91-5: Synthesized by immortalization of peripheral blood cells with Epstein-Barr virus. Gorny et al. [1989]

**No.** 50

MAb ID 1109/01

HXB2 Location p24 (69-86)

**Author Location** p24 (201–218 BRU)

Epitope LKETINEEAAEWDRVHPV

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG)

References Robert-Hebmann et al. 1992a; Robert-

Hebmann et al. 1992b

No. 51

MAb ID 14D4E11

HXB2 Location p24 (69-86)

Author Location p24 (201–218 BRU)

Epitope LKETINEEAAEWDRVHPV

Neutralizing no

Immunogen vaccine

Vector/Type: purified HIV-1

Species (Isotype) mouse (IgG1)

References Robert-Hebmann et al. 1992a; Robert-

Hebmann et al. 1992b; Janvier et al. 1992;

Janvier et al. 1990

 14D4E11: Mapped to aa209-217 through Pepscan method (original paper, AAEWDRVHP) – cross-reacts with HIV-2 Janvier *et al.* [1990] and to aa203-217 through EIA pentade-capeptide Janvier *et al.* [1992]. Janvier *et al.* [1990, 1992]

**No.** 52

MAb ID 1G5C8

**HXB2 Location** p24 (69–86)

**Author Location** p24 (201–218 BRU)

Epitope LKETINEEAAEWDRVHPV

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p24

Gag

Species (Isotype) mouse (IgG2b)

References Robert-Hebmann et al. 1992a; Robert-

Hebmann et al. 1992b; Janvier et al. 1992;

Janvier et al. 1990

1G5C8: Mapped to aa209-217 through Pepscan method (original paper, AAEWDRVHP) Janvier *et al.* [1990] and to aa203-217 through EIA pentadecapeptide Janvier *et al.* [1992]. Janvier *et al.* [1990, 1992]

**No.** 53

**MAb ID** 47-2

**HXB2 Location** p24 (69–86)

**Author Location** p24 (201–218 BRU)

Epitope LKETINEEAAEWDRVHPV

Neutralizing no

Immunogen vaccine

Strain: B clade BRU

Gag p24 Antibodies HIV Antibodies Tables

Species (Isotype) mouse (IgG) Epitope ETINEEAAEWD References Robert-Hebmann et al. 1992a: Robert-Neutralizing no Hebmann et al. 1992b Immunogen vaccine Vector/Type: beta-galactosidase fusion pro-No. 54 tein Strain: B clade IIIB HIV component: **MAb ID** 714/01 p24 Gag HXB2 Location p24 (69-86) Species (Isotype) mouse (IgG1) Author Location p24 (201-218 BRU) References Niedrig et al. 1991 Epitope LKETINEEAAEWDRVHPV • 113/038: cross-reactive between HIV-1, HIV-2 and SIV MAC Neutralizing no by multiple assays. Niedrig et al. [1991] Immunogen vaccine Strain: B clade IIIB HIV component: HIV-1 No. 58 Species (Isotype) mouse (IgG) **MAb ID** 1-E-4 References Robert-Hebmann et al. 1992a; Robert-**HXB2 Location** p24 (71–85) Hebmann et al. 1992b Author Location p24 (203-217) Epitope ETINEEAAEWDRVHP No. 55 Neutralizing no MAb ID polyclonal Immunogen vaccine HXB2 Location p24 (69-86) Strain: B clade IIIB HIV component: HIV-1 Author Location p24 (201–218 LAI) Species (Isotype) mouse (IgG) Epitope LKETINEEAAEWDRVHPV References Niedrig et al. 1989 Subtype B • 1-E-4: One of nine MAbs that bind to this peptide. Niedrig Neutralizing no et al. [1989] Immunogen vaccine Vector/Type: protein, virus-like particle (VLP) **No.** 59 Strain: B clade LAI HIV component: Gag, **MAb ID** 1-E-9 p17 Gag, p24 Gag Adjuvant: Complete HXB2 Location p24 (71-85) Freund's Adjuvant (CFA) **Author Location** p24 (203–217) Species (Isotype) mouse Epitope ETINEEAAEWDRVHP References Truong et al. 1997 **Neutralizing** no • An ELISA assay was used to study a panel of Gag peptides – Immunogen vaccine mature p24 CA epitopes mapped to residues 176-192, 201-218, Strain: B clade IIIB HIV component: HIV-1 233-253, 285-304, and were recognized by antibodies elicited Species (Isotype) mouse (IgG) by rp24CA - one p17MA epitope, residues 11-25, and one References Niedrig et al. 1989 p24CA epitope, residues 176-192, were recognized by anti-• 1-E-9: One of nine MAbs that bind to this peptide. Niedrig bodies raised against anti-p55 virus-like particles, suggesting et al. [1989] a different antigenic properties for p24CA and p17MA anti-No. 60 bodies depending on whether they are produced against the **MAb ID** 10-E-7 mature soluble protein or the immature assembled form of the HXB2 Location p24 (71-85) gag proteins. Truong et al. [1997] Author Location p24 (203–217) No. 56 Epitope ETINEEAAEWDRVHP MAb ID 111/073 Neutralizing no HXB2 Location p24 (71-81) Immunogen vaccine Author Location p24 (203-213 IIIB) Strain: B clade IIIB HIV component: HIV-1 Epitope ETINEEAAEWD Species (Isotype) mouse (IgG1) Neutralizing no References Niedrig et al. 1989; Niedrig et al. 1988 Immunogen vaccine • 10-E-7: One of nine MAbs that bind to this peptide - cross-Vector/Type: beta-galactosidase fusion proreactive with HIV-2 ROD and SIV MAC. Niedrig et al. [1989] tein Strain: B clade IIIB HIV component: • 10-E-7: Cross reactive between HIV-1, HIV-2 and SIV. Niedrig p24 Gag et al. [1988] Species (Isotype) mouse (IgG1) No. 61 References Niedrig et al. 1991 **MAb ID** 10-G-9 • 111/073: cross-reactive between HIV-1, HIV-2 and SIV MAC **HXB2 Location** p24 (71–85) by multiple assays. Niedrig et al. [1991] **Author Location** p24 (203–217) Epitope ETINEEAAEWDRVHP No. 57 **MAb ID** 113/038 Neutralizing no **HXB2 Location** p24 (71–81) Immunogen vaccine

Author Location p24 (203–213 IIIB)

Strain: B clade IIIB HIV component: HIV-1

**HIV Antibodies Tables** Gag p24 Antibodies

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1989: Niedrig et al. 1988

- 10-G-9: One of nine MAbs that bind to this peptide. Niedrig et al. [1989]
- 10-G-9: HIV-1 specific. Niedrig et al. [1988]

No. 62

MAb ID 11-C-5

HXB2 Location p24 (71-85)

Author Location p24 (203-217)

Epitope ETINEEAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1989; Niedrig et al. 1988

- 11-C-5: One of nine MAbs that bind to this peptide. Niedrig et al. [1989]
- 11-C-5: HIV-1 specific. Niedrig et al. [1988]

No. 63

MAb ID 2-E-4

**HXB2 Location** p24 (71–85)

Author Location p24 (203–217)

Epitope ETINEEAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG2a)

References Niedrig et al. 1989; Niedrig et al. 1988

- 2-E-4: One of nine MAbs that bind to this peptide crossreactive with HIV-2 ROD. Niedrig et al. [1989]
- ELISA, HIV-1 and HIV-2 by WB. Niedrig et al. [1988]

No. 64

**MAb ID** 2-H-4

HXB2 Location p24 (71-85)

Author Location p24 (203–217)

Epitope ETINEEAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1989; Niedrig et al. 1988

- 2-H-4: One of nine MAbs that bind to this peptide crossreactive with HIV-2 ROD. Niedrig et al. [1989]
- 2-H-4: Cross reactive between HIV-1, HIV-2 and SIV by ELISA, HIV-1 and HIV-2 by WB. Niedrig et al. [1988]

No. 65

**MAb ID** 8-D-2

HXB2 Location p24 (71-85)

**Author Location** p24 (203–217)

Epitope ETINEEAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG2a)

References Robert-Hebmann et al. 1992a; Robert-Hebmann et al. 1992b; Niedrig et al. 1989; Niedrig et al. 1988

- 8-D-2: One of nine MAbs that bind to this peptide. Niedrig et al. [1989]
- 8-D-2: HIV-1 specific. Niedrig et al. [1988]

No. 66

**MAb ID** 8-G-9

HXB2 Location p24 (71-85)

Author Location p24 (203-217)

Epitope ETINEEAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG)

References Niedrig et al. 1989

• 8-G-9: One of nine MAbs that bind to this peptide. Niedrig et al. [1989]

**No.** 67

**MAb ID** 8-H-7

**HXB2** Location p24 (71–85)

Author Location p24 (203–217)

Epitope ETINEEAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB HIV component: HIV-1

**Species (Isotype)** mouse (IgG3)

References Robert-Hebmann et al. 1992a; Robert-Hebmann et al. 1992b; Niedrig et al. 1989;

Niedrig et al. 1988

• 2-E-4: Cross reactive between HIV-1, HIV-2 and SIV by • 8-H-7: One of nine MAbs that bind to this peptide. Niedrig et al. [1989]

No. 68

**MAb ID** C5123

HXB2 Location p24 (71-85)

Author Location p24 (203-217 HXB2)

Epitope ETINEEAAEWDRVHP

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: viral lysate HIV component:

HIV-1

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

References Hinkula et al. 1990

• C5123: Epitope defined by peptide blocking of binding to native protein - WB reactive with p53 and p24. Hinkula et al. [1990]

No. 69

**MAb ID** 1-B-7

**HXB2 Location** p24 (76–85)

**Author Location** p24 (208–217 BH10)

Epitope EAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB

Species (Isotype) mouse (IgG1)

Gag p24 Antibodies HIV Antibodies Tables

References Niedrig et al. 1989; Niedrig et al. 1988

• 1-B-7: Reacts with two overlapping peptides, region of overlap is given – reacted with HIV-2 and SIV MAC. Niedrig *et al.* [1989]

**No.** 70

**MAb ID** 3-B-7

HXB2 Location p24 (76-85)

**Author Location** p24 (208–217 BH10)

Epitope EAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1989; Niedrig et al. 1988

• 3-B-7: Reacts with two overlapping peptides, region of overlap is given – reacted with HIV-2. Niedrig *et al.* [1989]

**No.** 71

MAb ID 6-D-12

HXB2 Location p24 (76-85)

**Author Location** p24 (208–217 BH10)

Epitope EAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1989; Niedrig et al. 1988

• 6-D-12: Reacts with two overlapping peptides, region of overlap is given – reacted with HIV-2. Niedrig *et al.* [1989]

**No.** 72

**MAb ID** 6-E-7

**HXB2 Location** p24 (76–85)

**Author Location** p24 (208–217 BH10)

Epitope EAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1989; Niedrig et al. 1988

• 6-E-7: Reacts with two overlapping peptides, region of overlap is given – reacted with HIV-2 and SIV MAC. Niedrig *et al.* [1989]

**No.** 73

MAb ID 8-D-5

**HXB2 Location** p24 (76–85)

**Author Location** p24 (208–217 BH10)

Epitope EAAEWDRVHP

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB

Species (Isotype) mouse (IgG)

References Niedrig et al. 1989; Niedrig et al. 1988

• 8-D-5: Reacts with two overlapping peptides, region of overlap is given – bound only HIV-1. Niedrig *et al.* [1989]

No. 74 MAb ID FF1 **HXB2 Location** p24 (76–90)

Author Location p24 (208–222 HXB2)

Epitope EAAEWDRVHPVHAGP

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: inactivated HIV

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Hinkula et al. 1990

 FF1: Epitope defined by peptide blocking of binding to native protein – WB reactive with p53 and p24. Hinkula et al. [1990]

**No.** 75

**MAb ID** 113/072

HXB2 Location p24 (81–90)

Author Location p24 (213–222 IIIB)

Epitope DRVHPVHAGP

Neutralizing no

Immunogen vaccine

Vector/Type: beta-galactosidase fusion protein Strain: B clade IIIB HIV component:

p24 Gag

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1991

• 113/072: Weak cross-reaction with HIV-2 on WB, otherwise not cross-reactive with HIV-2 or SIV MAC. Niedrig *et al.* [1991]

No. 76

**MAb ID** 25.3

**HXB2 Location** p24 (82–102)

Author Location p24 (82–102)

Epitope RVHPVHAGPIAPGQMREPRGS

Neutralizing no

Immunogen

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

References Momany et al. 1996

• 25.3: Crystal structure of the CA protein bound to Fab 25.3 was solved – monomers form 7 alpha-helices arranged in a coiled-coil – Fab binds to a long antigenic peptide that separates the longest helices, with a salt bridge at CA 82 R, and interactions as far away as positions 100 and 102. Momany *et al.* [1996]

**No.** 77

**MAb ID** 13-102-100

**HXB2 Location** p24 (84–94)

Author Location p24 (102–112 IIIB)

Epitope HPVHAGPIAPG

Neutralizing

Immunogen

Species (Isotype) mouse (IgG)

Research Contact Advanced Technologies, Inc., Columbia, MD References Qian & Tomer 1998; Parker *et al.* 1996

• 13-102-100: Affinity capillary electrophoresis was used to fine map this epitope, and the optimal peptide was defined as VHAGPIAPGIAP – this method uses migration time shifts to probe relative affinities of Abs – the antibody binds to the cyclophilin A binding domain. Qian & Tomer [1998]

HIV Antibodies Tables Gag p24 Antibodies

• 13-102-100: Binding site (HPVHAGPIAPG) defined by epitope footprinting – first binding p24 to MAb, then allowing proteolytic cleavage to take place to cleave unprotected residues, then performing mass spectrometry to identify protected residues of epitope. Parker *et al.* [1996]

**No.** 78

**MAb ID** RL4.72.1

**HXB2 Location** p24 (87–101)

Author Location p24 (219-233 BRU)

Epitope HAGPIAPGQMREPRG

Neutralizing no

Immunogen vaccine

Vector/Type: inactivated HIV Strain: D

clade NDK HIV component: HIV-1

Species (Isotype) mouse (IgG)

**References** Robert-Hebmann *et al.* 1992a; Robert-Hebmann *et al.* 1992b; Tatsumi *et al.* 1990

• RL4.72.1: Immunized with inactivated HIV NDK, D clade, reacts with B clade peptide. Robert-Hebmann *et al.* [1992a]

No. 79

**MAb ID** 406/01

**HXB2 Location** p24 (101–121)

Author Location p24 (233-253 BRU)

Epitope GSDIAGTTSTLQEQIGWMTNN

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB

Species (Isotype) mouse (IgG)

References Robert-Hebmann et al. 1992a; Robert-

Hebmann et al. 1992b

**No.** 80

MAb ID polyclonal

**HXB2 Location** p24 (101–121)

Author Location p24 (233–253 LAI)

Epitope GSDIAGTTSTLQEQIGWMTNL

Subtype B

Neutralizing no

Species (Isotype) mouse

Immunogen vaccine

Vector/Type: protein, virus-like particle (VLP) Strain: B clade LAI HIV component: Gag, p17 Gag, p24 Gag Adjuvant: Complete

Freund's Adjuvant (CFA)

References Truong et al. 1997

• An ELISA assay was used to study a panel of Gag peptides – mature p24 CA epitopes mapped to residues 176-192, 201-218, 233-253, 285-304, and were recognized by antibodies elicited by rp24CA – one p17MA epitope, residues 11-25, and one p24CA epitope, residues 176-192, were recognized by antibodies raised against anti-p55 virus-like particles, suggesting a different antigenic properties for p24CA and p17MA antibodies depending on whether they are produced against the mature soluble protein or the immature assembled form of the gag proteins. Truong et al. [1997]

**No.** 81

**MAb ID** 38:9.6K (38:96K)

**HXB2 Location** p24 (121–130)

Author Location p24 (253–262 HXB2)

Epitope NPPIPVGEIY

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p24-

p15 Gag

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Hinkula et al. 1990

- 38:9.6K: UK Medical Research Council AIDS reagent: ARP365.
- 38:9.6K: Called 38:96K epitope defined by peptide blocking of binding to native protein – WB reactive with p53 and p24. Hinkula *et al.* [1990]

**No.** 82

MAb ID EB1A9

HXB2 Location p24 (121-135)

**Author Location** p24 (253–267 LAI)

Epitope NPPIPVGEIYKRWII

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B

clade CBL-1 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

Research Contact R. B. Ferns and R. S. Tedder

References Ferns et al. 1989; Ferns et al. 1987

- EB1A9: UK Medical Research Council AIDS reagent: ARP345.
- EB1A9: Reacted with both p55 and p24 showed less than 75% homologous inhibition. Ferns *et al.* [1987]

**No.** 83

MAb ID polyclonal

**HXB2 Location** p24 (121–152)

Author Location Gag (253–284 LAI)

Epitope NPPIPVGEIYKRWIILGLNKIVRMYSPTSILD

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: p24 Gag Adjuvant: QS21

Species (Isotype) human (IgG)

References Pialoux et al. 2001

28 subjects were vaccinated with six HIV-1 peptides that were selected to be particularly rich in CTL epitopes, presented in lipopeptides with or without adjuvant QS21 – HIV-specific Ab responses were detected in 25/28 (89%), proliferative in 19/28 (79%), and CTL in 13/24 (54%) of testable volunteers – 25/28 had Ab responses to peptide G2, 14/28 had proliferative responses, and CTL responses were detected. Pialoux *et al.* [2001]

**No.** 84

MAb ID 30:3E5

**HXB2 Location** p24 (141–170)

Author Location p24 (273–302 HXB2)

Gag p24 Antibodies HIV Antibodies Tables

Epitope IVRMYSPTSILDIRQGPKEPFRDYVDRFYK

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: p24-

p15 Gag

**Species** (**Isotype**) mouse ( $IgG1\lambda$ )

Research Contact B. Wahren

References Hinkula et al. 1990

- 30:3E5: UK Medical Research Council AIDS reagent: ARP367.
- 30:3E5: Epitope defined by peptide blocking of binding to native protein – WB reactive with p53 and p24. Hinkula *et al.* [1990]

No. 85

MAb ID EF7

**HXB2 Location** p24 (141–170)

Author Location p24 (273–302 HXB2)

Epitope IVRMYSPTSILDIRQGPKEPFRDYVDRFYK

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: p24-

p15 Gag

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Lundin et al. 1996; Hinkula et al. 1990

- EF7: UK Medical Research Council AIDS reagent: ARP366.
- EF7: Included as a control. Lundin et al. [1996]
- EF7: Epitope defined by peptide blocking of binding to native protein WB reactive with p53. Hinkula *et al.* [1990]

No. 86

MAb ID LH-104-E

**HXB2 Location** p24 (143–148)

Author Location p24 (275–280 BRU)

Epitope RMYSPT

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade BRU

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Haaheim et al. 1991

- LH-104-E: UK Medical Research Council AIDS reagent: ARP319.
- LH-104-E: Reacts with both p24 and p55. Haaheim *et al.* [1991]

**No.** 87

**MAb ID** 1B2C12

**HXB2 Location** p24 (149–154)

Author Location p24 (273–292 IIIB)

**Epitope SILDIR** 

Neutralizing no

Immunogen vaccine

Vector/Type: purified HIV-1

Species (Isotype) mouse (IgG1)

**References** Janvier et al. 1992; Janvier et al. 1990

• 1B2C12: Reacts with HIV-1 and HIV-2— mapped to aa281-286 through Pepscan method Janvier *et al.* [1990], and to aa273-292 through EIA pentadecapeptide method Janvier *et al.* [1992]. Janvier *et al.* [1990, 1992]

No. 88

MAb ID LH-104-K

HXB2 Location p24 (149–154)

Author Location p24 (281-286 BRU)

Epitope SILDIR

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade BRU

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Haaheim et al. 1991

- LH-104-K: UK Medical Research Council AIDS reagent: ARP322.
- LH-104-K: Binds exclusively with p24 (not p55) Haaheim *et al.* [1991]

No. 89

MAb ID LH-104-A

**HXB2 Location** p24 (152–157)

**Author Location** p24 (BRU)

Epitope DIRQGP+QGVGGP

Neutralizing no

Immunogen vaccine

Vector/Type: peptide HIV component: p24

Gag

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

References Haaheim et al. 1991

- LH-104-A: UK Medical Research Council AIDS reagent: ARP307.
- LF-104-A: A 104 amino acid peptide was used to immunize mice hexapeptide scans revealed two reactive p24 peptides cross-competition studies indicated the region 270-286. Haaheim *et al.* [1991]

No. 90

**MAb ID** 1.17.3

**HXB2 Location** p24 (152–172)

**Author Location** p24 (152–172 SIVmac)

Epitope CVKQGPKEPFQSYVDRFYKSL

Neutralizing no

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade AGM TYO-7 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Otteken et al. 1992

• 1.17.3: Recognized an epitope present on HIV-2/SIVmac (MAC251/32H) and HIV-2smmH4, but not SIVagmTYO-1, HIV-1 IIIB or SIVmnd. Otteken *et al.* [1992]

**No.** 91

**MAb ID** 1A7

**HXB2 Location** p24 (152–172)

**Author Location** p24 (152–172 SIVmac)

Epitope CVKQGPKEPFQSYVDRFYKSL

Neutralizing no

Immunogen vaccine

HIV Antibodies Tables Gag p24 Antibodies

Vector/Type: inactivated HIV Strain: B clade AGM TYO-7 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Otteken et al. 1992

• 1A7: Recognized an epitope present on HIV-2/SIVmac (MAC251/32H) and HIV-2smmH4, but not SIVagmTYO-1, HIV-1 IIIB or SIVmnd. Otteken *et al.* [1992]

No. 92

MAb ID 1F6

**HXB2 Location** p24 (152–172)

Author Location p24 (152–172 SIVmac)

Epitope CVKQGPKEPFQSYVDRFYKSL

Neutralizing no Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade AGM TYO-7 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Otteken et al. 1992

• 1F6: Recognized an epitope present on HIV-2/SIVmac (MAC251/32H) and HIV-2smmH4, but not SIVagmTYO-1, HIV-1 IIIB or SIVmnd. Otteken *et al.* [1992]

No. 93

MAb ID 23A5G4

**HXB2 Location** p24 (153–172)

Author Location p24 (285–304 IIIB)

Epitope IRQGPKEPFRDYVDRFYKTL

Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: p24

Gag

Species (Isotype) mouse (IgG1)

**References** Janvier *et al.* 1996; Janvier *et al.* 1992; Janvier *et al.* 1990

- 23A5G4: A few sera which were able to bind the linear sequence 178-192, but not sequence 288-302 in an indirect peptide ELISA inhibited the binding of 23A5G4 to the native p24. Janvier *et al.* [1996]
- 23A5G4: Mapped to aa209-217 through Pepscan method Janvier *et al.* [1990] and to aa285-304 through EIA pentadecapeptide method Janvier *et al.* [1992]. Janvier *et al.* [1990, 1992]

**No.** 94

MAb ID 23A5G5

**HXB2 Location** p24 (153–172)

Author Location p24 (285–304 BRU)

Epitope IRQGPKEPFRDYVDRFYKTL

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: p24 Gag

Species (Isotype) mouse (IgG)

**References** Robert-Hebmann *et al.* 1992a; Robert-Hebmann *et al.* 1992b

No. 95

MAb ID 3D10G6

**HXB2 Location** p24 (153–172)

Author Location p24 (285–304 IIIB)

Epitope IRQGPKEPFRDYVDRFYKTL

Neutralizing no

Immunogen vaccine

Vector/Type: purified HIV-1

**Species (Isotype)** mouse (IgG1)

References Janvier et al. 1992; Janvier et al. 1990

 3D10G6: Epitope cross-reacts with HIV-1 and HIV-2- mapped to aa260-267 through Pepscan method Janvier et al. [1990] and to aa285-304 through EIA pentadecapeptide method. Janvier et al. [1990, 1992]

No. 96

MAb ID polyclonal

HXB2 Location p24 (153-172)

Author Location p24 (285–304 LAI)

Epitope IRQGPKEPFRDYVDRFYKTL

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein, virus-like particle (VLP) Strain: B clade LAI HIV component: Gag, p17 Gag, p24 Gag Adjuvant: Complete Freund's Adjuvant (CFA)

Species (Isotype) mouse

References Truong et al. 1997

• An ELISA assay was used to study a panel of Gag peptides – mature p24 CA epitopes mapped to residues 176-192, 201-218, 233-253, 285-304, and were recognized by antibodies elicited by rp24CA – one p17MA epitope, residues 11-25, and one p24CA epitope, residues 176-192, were recognized by antibodies raised against anti-p55 virus-like particles, suggesting a different antigenic properties for p24CA and p17MA antibodies depending on whether they are produced against the mature soluble protein or the immature assembled form of the gag proteins. Truong et al. [1997]

No. 97

MAb ID F5-4

HXB2 Location p24 (153-175)

Author Location p24 (153–174 HXB2)

Epitope IRQGPKEPFRDYVDRFYKTLRAE

Subtype B

Neutralizing no

Immunogen

Species (Isotype) mouse

References Kusk et al. 1992; Kusk et al. 1988

• F5-4: Binds to a location in the most hydrophilic region of p24. Kusk *et al.* [1988, 1992]

No. 98

**MAb ID** MO9.42.2

HXB2 Location p24 (153-178)

Author Location p24 (285–310 BRU)

Epitope IRQGPKEPFRDYVDRFYKTLRAEQAS

Neutralizing no

Immunogen vaccine

Vector/Type: virus Strain: HIV-2 ROD

HIV component: HIV-1

Species (Isotype) mouse (IgG)

Gag p24 Antibodies HIV Antibodies Tables

**References** Robert-Hebmann *et al.* 1992a; Robert-Hebmann *et al.* 1992b

MO9.42.2: Reacts with HIV-1s, HIV-2s, and SIVs in rec protein ELISA. Robert-Hebmann et al. [1992b]

No. 99

MAb ID MO9.50.2

**HXB2 Location** p24 (153–178)

Author Location p24 (285-310 BRU)

Epitope IRQGPKEPFRDYVDRFYKTLRAEQAS

Neutralizing no

Immunogen vaccine

Strain: HIV-2 ROD

Species (Isotype) mouse (IgG)

**References** Robert-Hebmann *et al.* 1992a; Robert-Hebmann *et al.* 1992b

• MO9.50.2: Reacts with HIV-1s, HIV-2s, and SIVs in rec protein ELISA. Robert-Hebmann *et al.* [1992b]

No. 100

MAb ID V10

**HXB2 Location** p24 (155–169)

**Author Location** p24 (289–303 IIIB)

Epitope QGPKEPFRDYVDRFY

Neutralizing no

Immunogen virus Species (Isotype) mouse

References Matsuo et al. 1992

 V10: Reacts with HIV-1 and SIV AGM analogous peptides. Matsuo et al. [1992]

**No.** 101

MAb ID V107

**HXB2 Location** p24 (155–177)

Author Location p24 (289-311 IIIB)

Epitope QGPKEPFRDYVDRFYKTLRAEQA

Neutralizing no

Immunogen virus

Species (Isotype) mouse

References Matsuo et al. 1992

V107: Reacts with FIV, HIV-1 and SIV AGM analogous peptides. Matsuo et al. [1992]

**No.** 102

MAb ID LH-104-C

**HXB2 Location** p24 (156–161)

**Author Location** p24 (BRU)

Epitope GPKEPF+QGVGGP

Neutralizing no

Immunogen vaccine

Vector/Type: peptide HIV component: p24

Gag

**Species (Isotype)** mouse (IgG3 $\kappa$ )

References Haaheim et al. 1991

- LH-104-C: UK Medical Research Council AIDS reagent: ARP309.
- LF-104-C: A 104 amino acid peptide was used to immunize mice hexapeptide scans revealed two reactive p24 peptides cross-competition studies indicated the region 351-373. Haaheim *et al.* [1991]

**No.** 103

MAb ID 12-B-4

HXB2 Location p24 (161-170)

Author Location p24 (293–302 IIIB)

Epitope FRDYVDRFYK

Neutralizing no

Immunogen vaccine

Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1989; Niedrig et al. 1988

• 12-B-4: Epitope is defined as the overlap between two HIV-1 reactive peptides – cross-reacts with HIV-2 ROD and SIV MAC. Niedrig *et al.* [1988, 1989]

**No.** 104

**MAb ID** C5122

HXB2 Location p24 (161-170)

Author Location p24 (293–302 HXB2)

Epitope FRDYVDRFYK

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: viral lysate HIV component:

HIV-1

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Hinkula et al. 1990

C5122: Defined by peptide blocking of binding to native protein – WB reactive with p53 and p24. Hinkula et al. [1990]

No. 105

MAb ID 9A4C4

HXB2 Location p24 (170–188)

Author Location p24 (303–317 IIIB)

Epitope KTLRAEQASQEVKNWMTET

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: p24 Gag

Species (Isotype) mouse (IgG1)

**References** Robert-Hebmann *et al.* 1992a; Robert-Hebmann *et al.* 1992b; Janvier *et al.* 1992;

Janvier et al. 1990

• 9A4C4: Mapped to aa260-267 through Pepscan method Janvier *et al.* [1990] – and to aa303-317 through EIA pentadecapeptide method Janvier *et al.* [1992]. Janvier *et al.* [1990, 1992]

**No.** 106

MAb ID 11C10B10

HXB2 Location p24 (171-185)

Author Location p24 (303–317 IIIB)

**Epitope** TLRAEQASQEVKNWM

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p24

Gao

Species (Isotype) mouse (IgG1)

References Janvier et al. 1992; Janvier et al. 1990

HIV Antibodies Tables Gag p24 Antibodies

• 11C10B10: Mapped to aa260-267 through Pepscan method Janvier *et al.* [1990] and to aa303-317 through EIA pentadecapeptide method Janvier *et al.* [1992]. Janvier *et al.* [1990, 1992]

**No.** 107

MAb ID 11D11F2

**HXB2 Location** p24 (171–185)

Author Location p24 (303-317 IIIB)

Epitope TLRAEQASQEVKNWM

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p24

Gag

Species (Isotype) mouse (IgG1)

References Janvier et al. 1992; Janvier et al. 1990

• 11D11F2: Mapped to aa260-267 through Pepscan method Janvier *et al.* [1990] and to aa303-317 through EIA pentadecapeptide method Janvier *et al.* [1992]. Janvier *et al.* [1990, 1992]

**No.** 108

MAb ID CD12B4

**HXB2 Location** p24 (171–185)

Author Location p24 (303–317 LAI)

Epitope TLRAEQASQEVKNWM

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B

clade CBL-1 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

Research Contact R. B. Ferns and R. S. Tedder

References Ferns et al. 1989; Ferns et al. 1987

- CD12B4: UK Medical Research Council AIDS reagent: ARP346.
- CD12B4: Reacted with both p55 and p24 strain-specific binding. Ferns et al. [1987]

**No.** 109

MAb ID BE3

**HXB2 Location** p24 (176–190)

Author Location p24 (308–322 HXB2)

Epitope QASQEVKNWMTETLL

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p24-

o15 Gag

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Research Contact B. Wahren

References Hinkula et al. 1990

- BE3: UK Medical Research Council AIDS reagent: ARP368.
- BE3: Defined by peptide blocking of binding to native protein
- WB reactive with p53 and p24. Hinkula et al. [1990]

**No.** 110

MAb ID L14

**HXB2 Location** p24 (176–190)

Author Location p24 (308–322 HXB2)

Epitope QASQEVKNWMTETLL

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p24-

p15 Gag

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Research Contact B. Wahren

References Hinkula et al. 1990

- L14: UK Medical Research Council AIDS reagent: ARP369.
- L14: Defined by peptide blocking of binding to native protein WB reactive with p53 and p24. Hinkula et al. [1990]

No. 111

**MAb ID** 108/03

HXB2 Location p24 (181-190)

**Author Location** p24 (313–322 IIIB)

**Epitope** VKNWMTETLL

Neutralizing no

Immunogen vaccine

Vector/Type: beta-galactosidase fusion protein Strain: B clade IIIB HIV component:

p24 Gag

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1991

• 108/03: Cross-reactive between HIV-1, HIV-2 and SIV MAC by multiple tests. Niedrig *et al.* [1991]

**No.** 112

MAb ID 110/015

**HXB2 Location** p24 (181–190)

Author Location p24 (313–322 IIIB)

Epitope VKNWMTETLL

Neutralizing no

Immunogen vaccine

Vector/Type: beta-galactosidase fusion protein Strain: B clade IIIB HIV component:

p24 Gag

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1991

• 110/015: Cross-reactive between HIV-1, HIV-2 and SIV MAC by multiple tests. Niedrig *et al.* [1991]

**No.** 113

**MAb ID** 32:32K

HXB2 Location p24 (199–222)

Author Location p24 (331–354 HXB2)

Epitope KTILKALGPAATLEEMMTACQGVG

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: p24-p15 Gag

**Species** (**Isotype**) mouse ( $IgG1\lambda$ )

References Hinkula et al. 1990

- 32:32K: UK Medical Research Council AIDS reagent: ARP368
- 32:32K: Epitope defined by peptide blocking of binding to native protein WB reactive with p53 and p24. Hinkula *et al.* [1990]

**No.** 114 **No.** 118 **MAb ID** C5200 MAb ID LH-104-B **HXB2 Location** p24 (199–222) **HXB2 Location** p24 (225–230) **Author Location** p24 (331–354 HXB2) Author Location p24 (357-362 BRU) Epitope KTILKALGPAATLEEMMTACQGVG **Epitope** GHKARV Subtype B Neutralizing no Immunogen vaccine **Neutralizing** Immunogen vaccine Vector/Type: peptide Strain: B clade BRU Vector/Type: viral lysate **Species (Isotype)** mouse (IgG1 $\kappa$ ) **Species (Isotype)** mouse (IgG1 $\kappa$ ) References Haaheim et al. 1991 References Hinkula et al. 1990 • LH-104-B: UK Medical Research Council AIDS reagent: • C5200: Epitope defined by peptide blocking of binding to ARP308. native protein. Hinkula et al. [1990] • LH-104-B: Binds exclusively with p55 (not p24), in contrast to LH-104-I. Haaheim et al. [1991] No. 115 MAb ID FH2 **No.** 119 **MAb ID** LH-104-I **HXB2 Location** p24 (201–215) Author Location p24 (333–347 HXB2) HXB2 Location p24 (226-231) **Epitope ILKALGPAATLEEMM** Author Location p24 (358–363 BRU) Subtype B **Epitope** HKARVL Neutralizing no Neutralizing no Immunogen vaccine Immunogen vaccine Vector/Type: protein HIV component: p24-Vector/Type: peptide Strain: B clade BRU **Species (Isotype)** mouse (IgG1 $\kappa$ ) p15 Gag **Species (Isotype)** mouse (IgG1 $\kappa$ ) References Haaheim et al. 1991 References Hinkula et al. 1990 • LH-104-I: UK Medical Research Council AIDS reagent: • FH2: Defined by peptide blocking of binding to native protein - WB reactive with p53 and p24. Hinkula et al. [1990] • LH-104-I: Binds exclusively with p24 (not p55), in contrast to LH-104-B. Haaheim et al. [1991] **No.** 116 **MAb ID** 13B5 **HXB2 Location** p24 (205–214) IV-C-4 Gag p24-p2p7p1p6 Antibodies Author Location p24 (205–213) **Epitope** LGPAATLEEM **No.** 120 **Neutralizing** MAb ID LH-104-G Immunogen vaccine **HXB2 Location** p24-p2p7p1p6 (231–5) Vector/Type: protein HIV component: p24 Author Location p24 (363–368 BRU) **Epitope LAEAMS** Species (Isotype) mouse Neutralizing no Ab Type C-term Immunogen vaccine Research Contact bioMerieux Vector/Type: peptide Strain: B clade BRU References Berthet-Colominas et al. 1999 **Species (Isotype)** mouse (IgG1 $\kappa$ ) • 13B5: Fab which was bound to p24 capsid for crystallization References Haaheim et al. 1991 and study of p24's structure. Berthet-Colominas et al. [1999] • LH-104-G: This epitope overlaps the p24-p2 cleavage site, database note. **No.** 117 • LH-104-G: UK Medical Research Council AIDS reagent: **MAb ID** 106/01 ARP320. HXB2 Location p24 (211-230) • LH-104-G: Reacts with both p24 and p55, in contrast to LH-Author Location p24 (343–362 IIIB) 104-I. Haaheim et al. [1991] Epitope LEEMMTACQGVGGPGHKARV Neutralizing no Immunogen vaccine Vector/Type: beta-galactosidase fusion protein Strain: B clade IIIB HIV component: p24 Gag Species (Isotype) mouse (IgG1) References Niedrig et al. 1991 • 106/01: Cross-reactive between HIV-1, HIV-2 and SIV MAC by multiple tests. Niedrig et al. [1991]

## IV-C-5 Gag p2p7p1p6 Antibodies

No. 121

MAb ID i5B11

HXB2 Location p2p7p1p6 (19-28)

Author Location p7 (5-14)

Epitope NFRNQRKIVK

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p7

Gag

Species (Isotype) rat (IgG2a)

References Tanchou et al. 1995; Tanchou et al. 1994; Otake et al. 1994

- i5B11: i5B11 and 15B11 may be two names for the same MAb.
- i5B11: MAb reacts with NCp7, NCp15, and partially inhibits NCp7-tRNA interaction. Tanchou et al. [1995]
- i5B11: Epitope mapped by ELISA and BIAcore inhibits NCp7 primer tRNA binding. Tanchou et al. [1994]

No. 122

MAb ID EC6

**HXB2** Location p2p7p1p6 (45–54) **Author Location** p15 (408–417 HXB2)

Epitope PRKKGCWKCG

Subtype B Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: p24-

p15 Gag

**Species (Isotype)** mouse (IgG2a $\kappa$ )

References Hinkula et al. 1990

• EC6: Epitope defined by peptide blocking of binding to native protein – WB reactive with p53. Hinkula et al. [1990]

**No.** 123

MAb ID M12

**HXB2** Location p2p7p1p6 (45–54) **Author Location** p15 (408–417 HXB2)

Epitope PRKKGCWKCG

Subtype B Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p24-

p15 Gag

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Hinkula et al. 1990

- M12: There is a p15 and a gp120 MAb both called M12.
- M12: Epitope defined by peptide blocking of binding to native protein – WB reactive with p53. Hinkula et al. [1990]

**No.** 124

MAb ID DG8

**HXB2 Location** p2p7p1p6 (66–81)

**Author Location** p7 (52–67)

Epitope RQANFLGKIWPSYKGR

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: p7 Species (Isotype) mouse (IgG)

Gag

Species (Isotype) mouse

References Tanchou et al. 1995

• DG8: Binds proximal to the second zinc-finger, inhibits NCp7tRNA interaction. Tanchou et al. [1995]

No. 125

MAb ID EB5

HXB2 Location p2p7p1p6 (66-81)

Author Location p7 (52-67)

Epitope RQANFLGKIWPSYKGR

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: p7

Species (Isotype) mouse

References Tanchou et al. 1995

• EB5: Binds proximal to the second zinc-finger – mutation at position 59 (Lys to Ser) results in 10-fold reduction in reactivity. Tanchou et al. [1995]

**No.** 126

MAb ID HH3

HXB2 Location p2p7p1p6 (66-81)

**Author Location** p7 (52–67)

Epitope RQANFLGKIWPSYKGR

Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: p7

Gag

**Species (Isotype)** mouse (IgG2b)

References Tanchou et al. 1995; Tanchou et al. 1994

- HH3: Binds proximal to the second zinc-finger. Tanchou et al. [1995]
- HH3: Epitopes mapped by ELISA and BIAcore does not inhibit NCp7 primer tRNA binding. Tanchou et al. [1994]

No. 127

MAb ID AD2

HXB2 Location p2p7p1p6 (78-86)

Author Location p7 (64-72)

Epitope YKGRPGNFL

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p7

Gag

Species (Isotype) mouse (IgG)

References Tanchou et al. 1995

• AD2: Binds at C term of NCp7. Tanchou et al. [1995]

**No.** 128

MAb ID CA5

HXB2 Location p2p7p1p6 (78-86)

**Author Location** p7 (64–72)

**Epitope** YKGRPGNFL

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p7

References Tanchou et al. 1995

Gag Antibodies HIV Antibodies Tables

• CA5: Binds at C term of NCp7. Tanchou et al. [1995] Neutralizing no Immunogen vaccine No. 129 Vector/Type: protein HIV component: p7 MAb ID DF3 Gag **HXB2 Location** p2p7p1p6 (78–86) Species (Isotype) mouse (IgG) **Author Location** p7 (64–72) References Tanchou et al. 1995 Epitope YKGRPGNFL • JB7: Binds at C term of NCp7. Tanchou et al. [1995] Neutralizing no Immunogen vaccine **No.** 134 MAb ID JF11 Vector/Type: protein HIV component: p7 HXB2 Location p2p7p1p6 (78-86) Gag Species (Isotype) mouse (IgG) **Author Location** p7 (64–72) References Tanchou et al. 1995 Epitope YKGRPGNFL • DF3: Binds at C term of NCp7. Tanchou et al. [1995] Neutralizing no Immunogen vaccine **No.** 130 Vector/Type: protein HIV component: p7 MAb ID EC3 Gag **HXB2 Location** p2p7p1p6 (78–86) Species (Isotype) mouse (IgG1) **Author Location** p7 (64–72) References Tanchou et al. 1995; Tanchou et al. 1994 Epitope YKGRPGNFL • JF11: Binds at C term of NCp7. Tanchou et al. [1995] Neutralizing no • JF11: Epitopes mapped by ELISA and BIAcore – does not Immunogen vaccine inhibit NCp7 primer tRNA binding. Tanchou et al. [1994] Vector/Type: protein HIV component: p7 Species (Isotype) mouse (IgG) **IV-C-6** Gag Antibodies References Tanchou et al. 1995 • EC3: Binds at C term of NCp7. Tanchou et al. [1995] **No.** 135 MAb ID polyclonal No. 131 HXB2 Location Gag (24-34) MAb ID FC12 **Author Location** Gag HXB2 Location p2p7p1p6 (78–86) Epitope GKTHYMINPL Author Location p7 (64-72) **Neutralizing** Epitope YKGRPGNFL Immunogen vaccine Neutralizing no Vector/Type: peptide HIV component: Env, Immunogen vaccine Gag, Nef, Pol Vector/Type: protein HIV component: p7 Species (Isotype) rabbit Gag References Li et al. 2005 Species (Isotype) mouse (IgG) Keywords mimics References Tanchou et al. 1995 • In early HIV-1 infection, patients develop autoimmune throm-• FC12: Binds at C term of NCp7, reacts with NCp15, inhibits bocytopenia, with Ab directed against beta3 integrin, GPIIIa49-NCp7-tRNA interaction. Tanchou et al. [1995] 66. Panning with a 7-mer phage display library using rabbit anti-GPIIIa49-66 (CAPESIEFPVSEARVLED), the immuno-**No.** 132 MAb ID GE4 dominant epitope of the identified potential molecular mimicry epitopes with HIV-1 Env (sklFDeGLFn, elfnkTIIFP), Pol **HXB2** Location p2p7p1p6 (78–86) (geAPEFPskq), Gag (gktHyMINPl) and Nef (qeeeeVgFPVt, **Author Location** p7 (64–72) qeeeeVgFPVt, edeGigFPVr, fkLVPVSEae, ssnTPTTNaa) pro-Epitope YKGRPGNFL teins. Pools of these peptides elicted Ab in rabbits that induce Neutralizing no platelet oxidation in vitro and thrombocytopenia in vivo upon Immunogen vaccine passive transfer. Nef (qeeeeVgFPVt), Gag (gktHyMINPl), and Vector/Type: protein HIV component: p7 Nef (fkLVPVSEae) all overlap with known HIV-1 epitopes. Li Gag et al. [2005] (mimics) **Species (Isotype)** mouse (IgG) References Tanchou et al. 1995 No. 136 • GE4: Binds at C term of NCp7. Tanchou et al. [1995] MAb ID 16/4/2 **HXB2 Location** Gag **No.** 133 Author Location p24 MAb ID JB7 **HXB2 Location** p2p7p1p6 (78–86) **Epitope Author Location** p7 (64–72) Neutralizing no

Immunogen vaccine

Epitope YKGRPGNFL

**HIV Antibodies Tables** Gag Antibodies

> Vector/Type: DNA with CMV promotor, DNA with CMV/MCK hybrid promotor, DNA with MCK promotor

Species (Isotype)

References Bojak et al. 2002a

• 16/4/2: The ability of three different promoters to induce Gag specific immune responses was compared. The cytomegaliovirus (CMV) early gene promoter, which allows constitutive expression in different cells of host tissue, the tissue spe- Research Contact A. O. Arthur, Frederick Cancer Research and cific muscle creatine kinase (MCK) promoter, which may be restricted to differentiated, multinucleated myofibers and so safer, and a hybrid MCK/CMV promoter – intramuscular immunization of BALB/c mice utilizing the MCK promoter in combination with a codon optimized gag gene generated humoral (IgG1 (Th1) and IgG2a (Th2)) and CTL immune responses against HIV-1 Gag, however, the quantified immune parameters were clearly reduced as compared to CMV promotor-driven Gag expression. Bojak et al. [2002a]

**No.** 137

MAb ID 183-H12-5C

**HXB2** Location Gag

Author Location p24

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype) mouse (IgG1)

Research Contact Bruce Chesebro and Kathy Wehrly, Rocky

Mountain Laboratories, Hamilton, Montana

References Wehrly & Chesebro 1997; Toohey et al. 1995; Chesebro et al. 1992

- 183-H12-5C: NIH AIDS Research and Reference Reagent Program: 3537.
- 183-H12-5C: Cross-reacts with HIV1 and HIV-2 p24, and SIV p27. Wehrly & Chesebro [1997]
- 183-H12-5C: Used as antigen capture reagent for p24 ELISA. Chesebro et al. [1992]; Toohey et al. [1995]

No. 138

**MAb ID** 241-D

**HXB2** Location Gag

Author Location p24

**Epitope** 

Neutralizing no

**Immunogen** 

**Species** (**Isotype**) human ( $IgG1\lambda$ )

Research Contact Susan Zolla-Pazner

las01@mcrcr6.med.nyu) (NYU Med.

Center)

**References** Robinson et al. 1991; Tyler et al. 1990; Gorny et al. 1989

- 241-D: MH AIDS Research and Reference Reagent program:
- 241-D: An antibody by this name is available in the NIH AIDS Research and Reference Reagent Program, and they refer to the papers Gorny et al. [1989]; Tyler et al. [1990]; Robinson et al. [1991], but no p24 MAb by this name is discussed in these papers. Gorny et al. [1989]; Robinson et al. [1991]; Tyler et al. [1990]

No. 139

MAb ID 2A6

**HXB2 Location** Gag

**Author Location** p17

**Epitope** 

Neutralizing

**Immunogen** 

Species (Isotype)

Development Center, Frederick, MD

References Pincus et al. 1998

2A6: Part of a panel of 17 MAbs used as controls testing for the dual specificity of MAb G11H3 for both p17 and mycoplasma. Pincus et al. [1998]

**No.** 140

MAb ID 5E2.A3k

**HXB2 Location** Gag

Author Location p24 (1-158 SF2)

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype) mouse (IgG1)

Research Contact Biodesign International, Kennebunk, Maine, **USA** 

References Hochleitner et al. 2000a

• 5E2.A3k: The Ab binding site was studied with epitope excision (protein is bound in native conformation to immobilized MAb, then digested with proteolytic enzymes) and extraction (protein is digested then allowed to react with Ab), followed by mass spectroscopy, as well as lysine modification - the epitope is discontinuous, but involves the highly conserved N-term proline, and the antibody recognizes SIVs and HIV-2 as well as HIV-1 p24. Hochleitner et al. [2000a]

**No.** 141

MAb ID 71-31

**HXB2 Location** Gag

**Author Location** p24

**Epitope** 

Neutralizing no

**Immunogen** 

 $(Z_0]$ -

**Species (Isotype)** human (IgG1 $\lambda$ )

References Bandres et al. 1998; Gorny et al. 1998; Gorny et al. 1997; Spear et al. 1993; Robinson et al. 1991; Robinson et al. 1990b; Gorny et al. 1989

- 71-31: NIH AIDS Research and Reference Reagent Program:
- 71-31: Included as a negative control in studies that demonstrate that CXCR4 can bind to gp120 in the absence of CD4gp120 interactions, and that this binding can be enhanced by Env deglycosylation. Bandres et al. [1998]
- 71-31: Did not mediate deposition of complement component C3 on HIV infected cells. Spear et al. [1993]
- 71-31: No enhancing or neutralizing activity. Robinson et al.
- 71-31: Did not enhance HIV-1 IIIB infection. Robinson et al. [1990b]

Gag Antibodies HIV Antibodies Tables

<b>No.</b> 142	• BC1071: The stoichiometry of MAb neutralization was tested
<b>MAb ID</b> 91-6	and MAb BC1071 was used in this study for virion quantifica-
HXB2 Location Gag	tion. Schonning et al. [1999]
Author Location p24 (121–240 IIIB)	No. 147
Epitope	No. 147 MAb ID BE10
Neutralizing no	
Immunogen HIV-1 infection	HXB2 Location Gag Author Location p7
Species (Isotype) human (IgG1 $\lambda$ )	Epitope
References Robinson et al. 1990b; Gorny et al. 1989	Neutralizing no
<ul> <li>91-6: NIH AIDS Research and Reference Reagent Program: 1239.</li> </ul>	Immunogen vaccine
• 91-6: No enhancing activity for HIV-1 IIIB. Robinson <i>et al.</i>	Vector/Type: protein HIV component: p7
[1990b]	Gag
[17700]	Species (Isotype) mouse (IgG)
<b>No.</b> 143	References Tanchou et al. 1995
<b>MAb ID</b> 98-4.3	• BE10: Binding NCp7 requires Zn fingers, does not react with
HXB2 Location Gag	NCp15, inhibits NCp7-tRNA interaction. Tanchou et al. [1995]
Author Location p24	NY 110
Epitope	No. 148
Neutralizing no	MAb ID CD9
Immunogen HIV-1 infection	HXB2 Location Gag
<b>Species</b> (Isotype) human (IgG1 $\lambda$ )	Author Location p7
<b>References</b> Robinson et al. 1991	Epitope
• 98-4.3: No enhancing or neutralizing activity. Robinson <i>et al.</i>	Neutralizing no
[1991]	Immunogen vaccine
<b>No.</b> 144	Vector/Type: protein HIV component: p7
MAb ID 98-4.9	Gag Species (Isotype) mouse (IgG)
HXB2 Location Gag	References Tanchou et al. 1995
Author Location p24	• CD9: Binds NCp7 independent of Zn fingers, does not react
Epitope	with NCp15. Tanchou <i>et al.</i> [1995]
Neutralizing no	with Nep13. Tanenou et al. [1993]
Immunogen HIV-1 infection	<b>No.</b> 149
Species (Isotype) mouse (IgG $3\lambda$ )	MAb ID CH9B2
<b>References</b> Gorny <i>et al.</i> 1989	HXB2 Location Gag
	Author Location p17
<b>No.</b> 145	Epitope
MAb ID AC2	Neutralizing
HXB2 Location Gag	Immunogen vaccine
Author Location p7	Vector/Type: inactivated HIV Strain: B
Epitope	clade CBL-1 HIV component: HIV-1
Neutralizing no	Species (Isotype) mouse (IgG1)
Immunogen vaccine	Research Contact R. B. Ferns and R. S. Tedder
Vector/Type: protein HIV component: p7	<b>References</b> Ferns et al. 1989; Ferns et al. 1987
Gag	• CH9B2: UK Medical Research Council AIDS reagent:
Species (Isotype) mouse (IgG)	ARP349.
References Tanchou et al. 1995	• CH9B2: Reactive against p18 and p55. Ferns et al. [1987]
• AC2: Binds NCp7 independent of Zn fingers, does not react	<b>No.</b> 150
with NCp15. Tanchou et al. [1995]	MAb ID ED8
<b>No.</b> 146	HXB2 Location Gag
MAb ID BC1071	Author Location p7
HXB2 Location Gag	Epitope
Author Location p24	Neutralizing no
Epitope	Immunogen vaccine
Neutralizing no	Vector/Type: protein HIV component: p7
Immunogen HIV-1 infection	Gag
Species (Isotype) mouse	Species (Isotype) mouse (IgG)
Research Contact Aalto BioReagents	References Tanchou et al. 1995
Deferences Schonning et al. 1000	

References Schonning et al. 1999

**HIV Antibodies Tables** Gag Antibodies

• ED8: Binds NCp7 independent of Zn fingers, does not react • HyHIV-19: Does not react with p17 peptides – Ka is 3.7 x 10<sup>6</sup> with NCp15. Tanchou et al. [1995]

No. 151

MAb ID EH12E1

**HXB2** Location Gag

Author Location p24

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade CBL-1 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

Research Contact R. B. Ferns and R. S. Tedder

References Ferns et al. 1989: Ferns et al. 1987

- EH12E1: UK Medical Research Council AIDS reagent: Research Contact R. B. Ferns and R. S. Tedder ARP313.
- EH12E1: Reacted with p55 and p24 in WB. Ferns et al. [1987]

No. 152

MAb ID G11G1

**HXB2 Location** Gag

Author Location p17

**Epitope** 

**Neutralizing** 

**Immunogen** 

Species (Isotype) rat

References Pincus et al. 1996; Shang et al. 1991

• G11G1: Immunotoxins were generated by linking Env MAbs to ricin A – immunotoxins mediated cell killing, but only if the antigen was expressed at the cell surface – ricin-G11G1 did not mediate cell killing. Pincus et al. [1996]

**No.** 153

MAb ID G11H3

**HXB2** Location Gag

Author Location p17

**Epitope** 

Neutralizing

Immunogen

Species (Isotype)

References Pincus et al. 1998; Shang et al. 1991

• G11H3: This MAb is cross-reactive between p17 and mycoplasma - this antibody binds strain specifically to the variable lipoprotein (Vlp) F of M. hyorhinis, in the region of the carboxy-terminal repeat CGGSTPTPEQGNNQGGSTPTPE-QGNSQVSK - the p17 epitope is discontinuous, but p17 and Vlp F share the tetrapeptide SQVS. Pincus et al. [1998]

**No.** 154

MAb ID HyHIV-19

**HXB2 Location** Gag

Author Location p17 (JMH1)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: p17

Gag

**Species (Isotype)** mouse (IgG1)

References Ota et al. 1998; Liu et al. 1995

M-1 for rec p17 – inhibited growth of HIV-1 JMH1 in MT-4 cells when added 24 hours after the initial culture. Ota et al. [1998]

No. 155

MAb ID IE8G2

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade CBL-1 HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Ferns et al. 1989; Ferns et al. 1987

- IE8G2: UK Medical Research Council AIDS reagent: ARP347.
- IE8G2: Reacted with both p55 and p24 broadly reactive - showed less than 75% homologous inhibition. Ferns et al.

No. 156

MAb ID V7-8

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** mouse (IgG3 $\kappa$ )

References Montefiori et al. 1991: Robinson et al. 1990b

- V7-8: NIH AIDS Research and Reference Reagent Program:
- V7-8: Reacted with HIV-1IIIB, RF, and MN. Montefiori et al.
- V7-8: Did not enhance HIV-1 IIIB infection. Robinson et al. [1990b]

No. 157

MAb ID anti-p24

**HXB2 Location** Gag

Author Location p24

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein, virus-like particle (VLP) HIV component: Gag, gp120, Nef, Pol

Species (Isotype) mouse (IgG)

Research Contact Intracel Co

References Buonaguro et al. 2001

• anti-p24: HIV-1 pr55 gag-based virus-like particles (VLP) carrying Nef and Pol open reading frames, as well as gp120 of the clade A isolate 94UG018, were created using a Baculovirus expression system to package additional ORFS into the VLP anti-V3 and anti-p24 Abs were used to assess the expression levels and Gag and gp120-TM were found to be expressed at comparable levels on the VLP. Buonaguro et al. [2001]

No. 158

Gag Antibodies HIV Antibodies Tables

MAb ID human sera

**HXB2 Location** Gag

Author Location p24

Epitope

Neutralizing

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG)

References Binley et al. 1997b

• Retention of anti-Env antibodies and loss of anti-Gag antibodies during progression was studied, and suggested to be the result of the loss of T-cell help and the unique ability of Env to stimulate B cells even in a backdrop of declining CD4 cells, because of the ability of Env to bind to the CD4 molecule. Binley *et al.* [1997b]

No. 159

MAb ID polyclonal

**HXB2 Location** Gag

Author Location Gag (LAI)

Epitope Subtype B Neutralizing

Immunogen vaccine

Vector/Type: DNA prime with protein boost Strain: B clade LAI HIV component: Gag,

Nef, Tat Adjuvant: IL-18

Species (Isotype) mouse

References Billaut-Mulot et al. 2001

 DNA vaccinated BALB/c mice primed and boosted with a multiepitopic vaccine with IL18 showed lymphoproliferative and CTL responses – co-administration of IL18 increased Tcell responses but decreased anti-HIV Ab levels. Billaut-Mulot et al. [2001]

**No.** 160

MAb ID polyclonal

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: gp120 depleted whole killed virus Strain: AG recombinant HZ321 HIV component: virus Adjuvant: Complete Freund's Adjuvant (CFA), CpG immunostimulatory sequence (ISS)

Species (Isotype) rat

References Moss et al. 2000

- Different HIV strains were used for different regions: subtype A env, subtype G gag.
- Lewis rats co-immunized with HIV-1 antigen in Freund's and with immunostimulatory sequences CpG stimulated increased IFN $\gamma$  expressing CD4+ and CD8+ T cells and anti-p24 anti-bodies relative to antigen in Freund's without CpG. Moss *et al.* [2000]

**No.** 161

MAb ID polyclonal

**HXB2 Location** Gag

**Author Location** p24 (SF2)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade SF2 HIV component: gp120, p24 Gag Adjuvant:

MF59, PLG

Species (Isotype) mouse

References O'Hagan et al. 2000

Microparticles were used as an adjuvant for entrapped HIV-1 gp120 and induced strong serum IgG responses in mice – polylactide co-glycolide polymer (PLG) microparticles in combination with MF-59 had the highest Ab response and also induced p24 specific CTL. O'Hagan et al. [2000]

**No.** 162

MAb ID polyclonal

**HXB2** Location Gag

**Author Location** Gag (SF2)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: DNA, protein Strain: B clade SF2 HIV component: Gag Adjuvant: aluminum phosphate, MF59, PLG

Species (Isotype) macaque, guinea pig, mouse

References O'Hagan et al. 2001

 DNA vaccines of codon-optimized Env and Gag genes driven by CMV promotors absorbed on to PLG microparticles were more effective than naked DNA at eliciting strong Ab responses (more rapid, higher titer, more stable), comparable to gp120 in MF-59. O'Hagan et al. [2001]

**No.** 163

MAb ID polyclonal

HXB2 Location Gag

**Author Location** p24

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HIV component: p24 Gag

Species (Isotype) rabbit (IgG)

References Gupta et al. 2001

 Gag p24 is the mostly widely used HIV protein for serological based diagnostic kits — phage display libraries of HIV-1 p24 identified 2 epitope-rich regions: 70% of the clones that were identified using immunized rabbit sera had DNA fragments from the N-terminal region spanning 150–240 of Gag, and 30% from the carboxy-terminal region of p24 containing amino acids 310–360 — subtype B and C comparisons were made. Gupta et al. [2001]

**No.** 164

MAb ID polyclonal

HXB2 Location Gag Author Location p55

Epitope

Neutralizing no

Immunogen vaccine

HIV Antibodies Tables Gag Antibodies

Vector/Type: protein, virus-like particle (VLP) Strain: B clade LAI HIV component: CD4BS, Gag, V3

Species (Isotype) mouse

References Truong et al. 1996

Antibodies raised against recombinant anti-p55 virus-like particles with the p24 region 196-226 deleted, bearing inserts of either the V3 or the CD4BS regions of gp120 were studied – no neutralizing responses, weak Env and strong Gag responses were elicited – the major homology region (MHR) and proximal sequences was found to be required for capsid assembly. Truong et al. [1996]

No. 165

MAb ID polyclonal

**HXB2** Location Gag

Author Location p24 (LAI)

Epitope Subtype B Neutralizing

Immunogen vaccine

Vector/Type: peptide, virion, baculovirus, E. Coli recombinant protein Strain: B clade LAI HIV component: p24 Gag Adjuvant: Complete Freund's Adjuvant (CFA), Incomplete Freund's Adjuvant (IFA)

Species (Isotype) rabbit (IgG)
References Devito et al. 2000c

• To compare vaccine strategies, rabbits were immunized with virion HIV-1/Lai, baculovirus recombinant p24, E. coli recombinant p24-15, and p24-derived peptides – the rabbit immunized with peptides had the broadest linear epitope responses – the capture ELISA method using anti-p24 IgG preparations was shown to capture isolates from HIV-1 subtypes or clades A to G – only immunization with virion HIV-1/Lai and baculovirus recombinant p24 developed IgG that was capable of efficiently capturing HIV-1 p24 in ELISA producing Abs able to recognise native configurations. Devito et al. [2000c]

No. 166

MAb ID polyclonal

**HXB2 Location** Gag

**Author Location** 

**Epitope Neutralizing** 

Immunogen vaccine

Vector/Type: DNA Adjuvant: CpG immunostimulatory sequence (ISS), phosphorothioate oligodeoxynucleotides (ODNs)

Species (Isotype) mouse

References Deml et al. 2001

Immunization mice with a codon-optimized Gag was compared with a non-optimized Rev dependent Gag expression vector

 Gag expression was at higher levels and Rev independent with the codon-optimized Gag, and i.m. immunization gave a stronger Th1-driven humoral and cellular immune response – intradermal immunization with either Gag DNA induced a Th2 response and no CTL. Deml et al. [2001]

**No.** 167

MAb ID polyclonal

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

**Neutralizing** yes

Immunogen HIV-1 infection

Species (Isotype) human

References Montefiori et al. 2001

• In 7/9 patients in whom HAART was initiated during early seroconversion, NAbs to autologous strains were not found immediately following treatment interuption after 1-3 years, and Env and Gag Abs were low or undetected by ELISA indicating, that early HAART suppresses the normal antibody response to HIV-1, presumably by limiting the concentration of viral antigens needed to drive B-cell maturation – in 3 patients with a viral rebound autologous NAbs rapidly appeared and correlated with spontaneous down-regulation of viremia - prolonged control of viremia after stopping treatment persisted in the absence of detectable NAbs, suggesting that cellular immune responses alone can control viremia under certain circumstances - these results support the notion that virus-specific B-cell priming, combined with CD8+ CTL induction, may be beneficial for HIV-1 vaccines that aim to suppress viremia in the absence of complete protection to prevent disease and reduce the rate of virus transmission. Montefiori et al. [2001]

**No.** 168

MAb ID polyclonal

**HXB2 Location** Gag

**Author Location** 

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: virus-like particle (VLP) HIV component: Env, Gag Adjuvant: Complete Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgG)

References Lebedev et al. 2000

 Virus-like particles (VLPs) in the form of spherical particles with yeast dsRNA enveloped in a polysaccharide matrix carrying the protein TBI, that contains fragments of HIV Env and Gag, were used to immunize BALB/c mice and induced specific Abs against HIV-1 as measured by ELISA with TBI. Lebedev et al. [2000]

**No.** 169

MAb ID polyclonal

**HXB2** Location Gag

**Author Location** 

**Epitope** 

Neutralizing no

Immunogen vaccine

*Vector/Type:* DNA with CMV promotor, DNA with CMV/MCK hybrid promotor, DNA with

MCK promotor

Species (Isotype) mouse (IgG1, IgG2a)

References Bojak et al. 2002a

Gag Antibodies HIV Antibodies Tables

• The ability of three different promoters to induce Gag specific immune responses was compared. The cytomegaliovirus (CMV) early gene promoter, which allows constitutive expression in different cells of host tissue, the tissue specific muscle creatine kinase (MCK) promoter, which may be restricted to differentiated, multinucleated myofibers and so safer, and a hybrid MCK/CMV promoter – intramuscular immunization of BALB/c mice utilizing the MCK promoter in combination with a codon optimized gag gene generated humoral (IgG1 (Th1) and IgG2a (Th2)) and CTL immune responses against HIV-1 Gag, however, the quantified immune parameters were clearly reduced as compared to CMV promotor-driven Gag expression. Bojak *et al.* [2002a]

**No.** 170

MAb ID polyclonal

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human

References Meles et al. 2002

• Indeterminant WB in Ethiopians: of 12,124 specimens blood specimens from Ethiopia, 1,437 (11.9%) were HIV-1-positive for antibody, and 91 (0.8%) gave equivocal results, most often due to p24 reactivity – subsequent testing confirmed many of the indeterminants were HIV-negative – the American Red Cross diagnostic criteria was more accurate than CDC or WHO, which would have given some false positive results. Meles *et al.* [2002]

**No.** 171

MAb ID polyclonal

**HXB2 Location** Gag

Author Location p24

**Epitope** 

Subtype A

Neutralizing yes

Immunogen vaccine

Vector/Type: virus-like particle (VLP) Strain: A clade UG5.94UG018 HIV com-

ponent: Gag, gp120

Species (Isotype) mouse

References Buonaguro et al. 2002

Country Uganda

**Keywords** subtype comparisons

• BALB/c mice were immunized with VLPs carrying a subtype A gp120. Humoral immune responses directed against B-clade derived Gag (p24) peptides or gp120-Env V3 loop peptide were readily induced following a multi-dose immunization with VLP particles presenting a gp120 molecule from a HIV-1 isolate of clade A. VLP-immunized mice showed autologous and heterologous (against B-clade HIV-1 IIIB strain) neutralization activity. Proliferative responses and CTL were also observed. Buonaguro et al. [2002] (subtype comparisons)

**No.** 172

MAb ID polyclonal

**HXB2 Location** Gag

**Author Location** Gag

Epitope

**Neutralizing** 

Immunogen vaccine

Vector/Type: DNA HIV component: Gag

Species (Isotype) mouse (IgG1)

References Bojak et al. 2002b

Keywords Th1

• Balb/c mice vaccinated by syngag, a DNA plasmid expressing HIV-1 Gag modified for human/mammalian codon usage, gave stronger and longer lasting immune responses than wild type gag. Gag-specific antibody and cellular immune responses were both increased, with a clear T-helper 1 polarization. There was a better IgG1/IgG2 response to intramuscular (i.m.) as compared to subcutaneous (s.c.) vaccination. Bojak *et al.* [2002b] (**Th1**)

**No.** 173

MAb ID polyclonal

**HXB2 Location** Gag

**Author Location** Gag

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: DNA, protein, virus-like particle (VLP), PLG microparticle Adjuvant: E. coli heat labile enterotoxin

Species (Isotype) macaque

References Otten et al. 2003

• This study evaluates different vaccine technologies that avoid live vectors including plasmid DNA, recombinant p55Gag protein or gag-pol administered by polylactide coglycolide (PLG) microparticles, LTK63 as adjuvant, VLP, and plasmid DNA. 4/4 macaques primed with Gag-PLG and LTK63 showed strong antibody responses after the fourth immunization at week six. The best CTL responses were found for gag DNA, the best Th and Ab were obtained using Gag protein on PLG microparticles; Gag DNA priming with a PLG-protein boost gave high level CTL, Th and Ab responses. Otten *et al.* [2003]

**No.** 174

MAb ID polyclonal

**HXB2** Location Gag

Author Location p24

**Epitope** 

Subtype multiple

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

References Barin et al. 2005

Country France

Keywords acute/early infection, assay development

A combination of 4 antigenic regions was used to differentiate between early (<180 days) and chronic infection. These regions were: p24; the gp41 peptide spanning the immunodominant epitope (IDE) of gp41, RVAVERYLKDQQLLGIWGC-SGKICTTAV, and a subtype D version of this peptide; 5 V3 consensus peptides including A, B, C, D, and CRF01-AE; and Integrase. V3 and the IDE provide the best discrimination, with >20 fold higher levels in chronic infection when assayed by

HIV Antibodies Tables Protease Antibodies

EIA using dried serum spots. Antibodies to Integrase and p24 were not as distinctive, and people tend to lose, not increase, responses to p24 over time. This assay can be used to identify samples from early infection with high sensitivity and specificity. Barin *et al.* [2005] (assay development, acute/early infection)

No. 175

MAb ID polyclonal HIVIG

**HXB2 Location** Gag **Author Location** p24

Epitope

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Nichols et al. 2002

 NYBC-HIVIG derived from patients with high NAb titers and NABI-HIVIG derived from patients with high anti-p24 Ab titers were compared in neutralizing assay against a panel of six primary isolates—both could neutralize all isolates tested but the NYBC-HIVIG dose required for 50% neutralization was of 3.2 fold lower, showing that the source plasmas influence the effective concentration of NAb present in HIVIG. Nichols et al. [2002]

#### **IV-C-7** Protease Antibodies

**No.** 176

**MAb ID** 1696

HXB2 Location Protease (1-7)

**Author Location** Protease (1–7 BH10)

Epitope PQIYLWQ

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: Pro-

tease

Species (Isotype) mouse (IgG)

Ab Type N-term

References Lescar et al. 2003; Rezacova et al. 2002; Reza-

cova et al. 2001; Lescar et al. 1999

Keywords review, structure

- 1696: Study compares the crystal structure of the scFv-1696 in the non-complexed form compared to the complexed Fab-1696 and the Ag-bound scFv-1696 structures. Changes in the three conformational tertiary structures of CDR-H3 as well as in the different relative orientations of the light-chain variable domains of the different structures were observed, demonstrating plasticity in the antibody binding site. Lescar *et al.* [2003] (structure)
- 1696: Review of the implications of antibody structure and antigen peptide binding for the mechanisms of inhibition of protease activity by two MABs with different binding sites in protease. Rezacova *et al.* [2002] (**review**, **structure**)
- 1696: The crystal structure of the single chain Fv fragment of 1696 bound to a cross-reactive peptide (PQITLWQRR) was obtained. This sturcture suggests that 1696 inhibits portease activity by favoring dissociation of the active homodimer. Rezacova *et al.* [2001] (**structure**)

• 1696: MAb binds to HIV-1 and HIV-2, putative epitopes are PQIYLWQ and PQFSLWK respectively – Pro1 is critical, QIYLWQR residues 2-8, does not compete - MAb disrupts catalytic activity – crystal structure of the ligand-free Fab at 3 A resolution reveals a deep cavity lined by acidic and hydrophobic residues – the binding region is located within the region required for dimerization and the Fab structure could serve as a basis for drug design targeting this region. Lescar et al. [1999] (structure)

**No.** 177

**MAb ID** 10E7

HXB2 Location Protease (36–46)

**Author Location** Protease (38–45 HXB2)

Epitope MSLPGRWKPKM

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Pro-

tease

Species (Isotype) hamster (IgG)

References Bjorling et al. 1992; Croix et al. 1993

10E7: Immunodominant region of protease in Armenian hamster (but only weakly reactive in people, see: Bjorling *et al.* [1992]) – peptide MSLPGRWKP blocks protease binding Croix *et al.* [1993]. Bjorling *et al.* [1992]; Croix *et al.* [1993]

**No.** 178

**MAb ID** F11.2.32

**HXB2 Location** Protease (36–46)

**Author Location** Protease (36–46 BH10)

Epitope MSLPGRWKPKM

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: Protease

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type flap region

References Rezacova et al. 2002; Lescar et al. 1999;

Lescar et al. 1997: Lescar et al. 1996

**Keywords** review, structure

- F11.2.32: Review of the implications of antibody structure and antigen peptide binding for the mechanisms of inhibition of protease activity by two MABs with different binding sites in protease. Rezacova *et al.* [2002] (**review**, **structure**)
- F11.2.32: Crystal structure of a Fab peptide complex was obtained. Distortion may occur in the flap region of the protein, important for regulating access of substrate to the catalytic site. Lescar *et al.* [1999] (**structure**)
- F11.2.32: Binding leads to significant inhibition in proteolytic activity crystal structure of Fab-peptide was determined to 2.2 A resolution bound peptide shows no structural similarity to the corresponding segment in native protease suggesting binding may distort protein structure. Lescar *et al.* [1997] (structure)

**No.** 179

MAb ID 13E1

HXB2 Location Protease (38–45)

Author Location Protease (38–45 HXB2)

RT Antibodies HIV Antibodies Tables

**Epitope** LPGRWKPK Subtype B Neutralizing no Immunogen vaccine Vector/Type: protein HIV component: Pro-Species (Isotype) hamster (IgG) References Croix et al. 1993 • 13E1: Binds to MSLPGRWKPKM with sightly higher affinity. Croix et al. [1993] **No.** 180 MAb ID 8B11 **HXB2 Location** Protease (38–45) **Author Location** Protease (38–45 HXB2) Epitope LPGRWKPK Subtype B Neutralizing no Immunogen vaccine Vector/Type: protein HIV component: Protease Species (Isotype) hamster (IgG) References Croix et al. 1993 • 8B11: Binds to MSLPGRWKPKM with sightly higher affinity. Croix et al. [1993] No. 181 **MAb ID** 8C10 **HXB2 Location** Protease (38–45) **Author Location** Protease (38–45 HXB2) **Epitope** LPGRWKPK Subtype B Neutralizing no Immunogen vaccine Vector/Type: protein HIV component: Protease Species (Isotype) hamster (IgG) References Croix et al. 1993 • 8C10: Binds to MSLPGRWKPKM with sightly higher affinity. Croix et al. [1993] **No.** 182 MAb ID 8G5 **HXB2 Location** Protease (38–45) **Author Location** Protease (38–45 HXB2) Epitope LPGRWKPK Subtype B Neutralizing no

Vector/Type: protein HIV component: Pro-

• 8G5: Binds to MSLPGRWKPKM with sightly higher affinity.

#### **IV-C-8** RT Antibodies

**No.** 183

MAb ID 1E8

HXB2 Location RT (65–73)

**Author Location** RT (65–73)

Epitope KKDSTKWRK

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: RT

Adjuvant: nitrocellulose

Species (Isotype) mouse (IgG1)

References Gu et al. 1996; Wu et al. 1993

- 1E8: Significantly inhibits DNA polymerase activity of RT by hindering binding of dNTPs additive or synergistic RT inhibition with nevirapine and delayirdine. Gu *et al.* [1996]
- 1E8: Inhibits RT activity, binding site overlaps with two AZT resistance mutations. Wu et al. [1993]

No. 184

MAb ID polyclonal

**HXB2 Location** RT (249-263)

Author Location RT (249–263)

Epitope KDSWTVNDIQKLVGK

**Neutralizing** 

Immunogen vaccine, in vitro stimulation or selection

Vector/Type: peptide presented on icosahedral protein scaffold HIV component: RT Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (Isotype) human (IgG)

Ab Type gp120 C2

References Domingo et al. 2003

Keywords vaccine antigen design

• A virus-like protein scaffold, called E2DISP, derived from pyruvate dehydrogenase multienzyme from Bacillus stearothermophilus has been engineered to display 60 copies of one or more epitopes on a single molecule. An E2DISP scaffold which displayed pep23, a 15-residue B and T helper HIV-1 RT epitope elicited a pep23-specific T-helper response in vitro. The E2DISP scaffold displaying peptide RT2, which is a CTL HIV-1 RT epitope, was able to elicit a CD8+ T cell response in vitro and in a vaccinated HLA-A2 transgenic mouse. Thus the E2DISP scaffold allows cell-entry and access to both the class I and class II processing pathways. The Th response in vaccinated mice supported Pep23-specific IgG responses. Domingo et al. [2003] (vaccine antigen design)

**No.** 185

**MAb ID** 1.152 B3

**HXB2 Location** RT (294–302)

Author Location RT (294-302)

**Epitope PLTEEAELE** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: RT

Species (Isotype) mouse (IgG1)

References Orvell et al. 1991

• 1.152 B3: Weakly positive by immunofluorescence – binding inhibits RT enzymatic activity. Orvell *et al.* [1991]

Immunogen vaccine

Species (Isotype) hamster (IgG)

Croix et al. [1993]

tease

References Croix et al. 1993

**HIV Antibodies Tables** RT Antibodies **No.** 186 Neutralizing no **MAb ID** 1.158 E2 Immunogen vaccine **HXB2 Location** RT (294–302) Vector/Type: E. coli Trp fusion protein HIV Author Location RT (294-302) component: RT Species (Isotype) mouse (IgG1) **Epitope PLTEEAELE** Neutralizing no References Szilvay et al. 1992 Immunogen vaccine • 33D5: Weak inhibitor of RT, reactive by immunofluorescence. Vector/Type: protein HIV component: RT Szilvay et al. [1992] Species (Isotype) mouse (IgG1) No. 191 References Orvell et al. 1991 MAb ID 5B2 • 1.158 E2: Negative by immunofluorescence – binding inhibits **HXB2 Location** RT (294–318) RT enzymatic activity. Orvell et al. [1991] Author Location RT (294-319) **No.** 187 Epitope PLTEEAELELAENREILKEPVHGVY **MAb ID** 31D6 Neutralizing no **HXB2 Location** RT (294–318) Immunogen vaccine Author Location RT (294-319) Vector/Type: E. coli Trp fusion protein HIV Epitope PLTEEAELELAENREILKEPVHGVY component: RT Neutralizing no Species (Isotype) mouse (IgG1) Immunogen vaccine References Szilvay et al. 1992 *Vector/Type:* E. coli Trp fusion protein HIV • 5B2: UK Medical Research Council AIDS reagent: ARP3018. component: RT • 5B2: There is an RT specific Ab Szilvay et al. [1992] and a Species (Isotype) mouse (IgG1) gp41 specific Ab Tian et al. [2001] both called 5B2. Szilvay References Szilvay et al. 1992 et al. [1992] • 31D6: Strong inhibitor of RT, > 50% inhibition. Szilvay et al. • 5B2: Weak inhibitor of RT, reactive by immunofluorescence. [1992] Szilvay et al. [1992] No. 188 No. 192 **MAb ID** 31G8 MAb ID polyclonal **HXB2 Location** RT (294–318) HXB2 Location RT (295-304) **Author Location** RT (294–319) **Author Location** RT (295–304 PV22) Epitope PLTEEAELELAENREILKEPVHGVY Epitope LTEEAELELA Neutralizing no Neutralizing no Immunogen vaccine Immunogen HIV-1 infection Vector/Type: E. coli Trp fusion protein HIV Species (Isotype) human (IgG) References Grimison & Laurence 1995 component: RT Species (Isotype) mouse (IgG1) No. 193 References Szilvay et al. 1992 MAb ID 1.153 G10 • 31G8: Weak inhibitor of RT, reactive by immunofluorescence. HXB2 Location RT (350-354) Szilvay et al. [1992] Author Location RT (350-354) **No.** 189 Epitope KTGKY MAb ID 32E7 Neutralizing no **HXB2 Location** RT (294-318) Immunogen vaccine Author Location RT (294-319) Vector/Type: protein HIV component: RT Epitope PLTEEAELELAENREILKEPVHGVY Species (Isotype) mouse (IgG1) Neutralizing no References Orvell et al. 1991

Immunogen vaccine Vector/Type: E. coli Trp fusion protein HIV component: RT Species (Isotype) mouse (IgG1)

• 32E7: Weak inhibitor of RT, reactive by immunofluorescence. Szilvay et al. [1992]

No. 190 **MAb ID** 33D5 **HXB2 Location** RT (294–318) **Author Location** RT (294–319)

References Szilvay et al. 1992

Epitope PLTEEAELELAENREILKEPVHGVY

No. 194 MAb ID RTMAb8 **HXB2 Location** RT (376-383) **Author Location** RT (532–539) **Epitope TTESIVIW** Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: RT

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Species (Isotype) mouse (IgG)

References Ferns et al. 1991; Tisdale et al. 1988

No. 195

RT Antibodies HIV Antibodies Tables

MAb ID 1D4A3 **Epitope** Subtype B **HXB2 Location** RT (384–387) Author Location RT (540-543) Neutralizing Epitope GKIP Immunogen HIV-1 infection Neutralizing no Species (Isotype) human Immunogen vaccine Research Contact Amon Hizi, Sackler School of Medicine, Tel Vector/Type: protein HIV component: RT Aviv, Isreal Species (Isotype) mouse (IgG) References Herschhorn et al. 2003 Keywords antibody generation, antibody sequence, vari-References Ferns et al. 1991 able domain, immunotherapy **No.** 196 • 5B11: One of five human single chain Fv (ScFv) Abs isolated MAb ID RT6H from an phage display library. F-6 was shown to the carboxyl **HXB2 Location** RT (384–387) terminal segment of the p66 RT polypeptide, corresponding to Author Location RT (540-543) RNase H. F-6 inhibited the DNA and RNA-dependent DNA **Epitope** GKIP polymerase (RDDP) activities of HIV-1 RT; two others, 6E9 Neutralizing no and 5B11, inhibited RDDP and so have possible therapeutic Immunogen vaccine potential. Herschhorn et al. [2003] (antibody generation, im-Vector/Type: protein HIV component: RT munotherapy, antibody sequence, variable domain) Species (Isotype) mouse (IgG) References Ferns et al. 1991 No. 201 **MAb ID** 6B10 **No.** 197 **HXB2 Location RT** MAb ID 1.160 B3 Author Location RT (BH-10) **HXB2 Location** RT (442–450) **Epitope** Author Location RT (442–450) Subtype B **Epitope** VDGAANRET **Neutralizing** Neutralizing no Immunogen HIV-1 infection Immunogen vaccine Species (Isotype) human Vector/Type: protein HIV component: RT Research Contact Amon Hizi, Sackler School of Medicine, Tel Species (Isotype) mouse (IgG1) Aviv, Isreal References Orvell et al. 1991 References Herschhorn et al. 2003 Keywords antibody generation, antibody sequence, vari-**No.** 198 able domain MAb ID polyclonal • 6B10:One of five human single chain Fv (ScFv) Abs isolated **HXB2 Location** RT (521–531) from an phage display library. F-6 was shown to the carboxyl **Author Location** RT (521–531 PV22) terminal segment of the p66 RT polypeptide, corresponding to Epitope IIEQLIKKEKV RNase H. F-6 inhibited the DNA and RNA-dependent DNA Neutralizing no polymerase (DDDP and RDDP) activities of HIV-1 RT; two Immunogen HIV-1 infection others, 6E9 and 5B11, inhibited RDDP and so have possible Species (Isotype) human (IgG) therapeutic potential. In contrast, 6B10 seemed to enhance References Grimison & Laurence 1995 DDDP activity and did not effect RDDP. Herschhorn et al. [2003] (antibody generation, antibody sequence, variable No. 199 domain) **MAb ID** C2003 **HXB2 Location** RT (536–549) No. 202 **Author Location** RT (703–716 BH10) MAb ID 6E9 Epitope VPAHKGIGGNEQVD **HXB2 Location RT** Neutralizing no Author Location RT (BH-10) Immunogen vaccine **Epitope** Vector/Type: peptide Strain: B clade BH10 Subtype B Species (Isotype) rabbit (IgG) **Neutralizing** References DeVico et al. 1991 Immunogen in vitro stimulation or selection • C2003: Inhibits polymerase activity from a variety of retro-Species (Isotype) human viruses – RT protected from inhibition by preincubation with Research Contact Amon Hizi, Sackler School of Medicine, Tel template primer. DeVico et al. [1991] Aviv, Isreal References Herschhorn et al. 2003 **No.** 200 Keywords antibody generation, antibody sequence, vari-MAb ID 5B11

**HXB2 Location** RT **Author Location** RT (BH-10)

able domain, immunotherapy

HIV Antibodies Tables RT Antibodies

• 6E9: One of five human single chain Fv (ScFv) Abs isolated from an phage display library. F-6 was shown to the carboxyl terminal segment of the p66 RT polypeptide, corresponding to RNase H. F-6 inhibited the DNA and RNA-dependent DNA polymerase (RDDP) activities of HIV-1 RT; two others, 6E9 and 5B11, inhibited RDDP and so have possible therapeutic potential. Herschhorn *et al.* [2003] (antibody generation, immunotherapy, antibody sequence, variable domain)

No. 203 MAb ID E-4 HXB2 Location RT

Author Location RT (BH-10)

Epitope Subtype B Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Research Contact Amon Hizi, Sackler School of Medicine, Tel Aviv, Isreal

References Herschhorn et al. 2003

**Keywords** antibody generation, antibody sequence, variable domain

• E-4:One of five human single chain Fv (ScFv) Abs isolated from an phage display library. F-6 was shown to the carboxyl terminal segment of the p66 RT polypeptide, corresponding to RNase H. F-6 inhibited the DNA and RNA-dependent DNA polymerase (RDDP) activities of HIV-1 RT; two others, 6E9 and 5B11, inhibited RDDP and so have possible therapeutic potential. In contrast, E-4 seemed to enhance RDDP. Herschhorn *et al.* [2003] (antibody generation, antibody sequence, variable domain)

**No.** 204

MAb ID 6B9

**HXB2 Location RT** 

**Author Location** RT (155–250)

Epitope Neutralizing yes Immunogen vaccine

Vector/Type: vaccinia Strain: B clade HXB2 HIV component: RT

Species (Isotype) mouse (IgG)

**Ab Type** RT palm domain

**References** Ohba *et al.* 2001; Chiba *et al.* 1997; Chiba *et al.* 1996

• 6B9: In contrast to MAb 7C4, which binds to the thumb region of RT, 6B9 binds to the palm subdomain and does not inhibit RT activity. Chiba *et al.* [1996]

No. 205

MAb ID 5F

**HXB2 Location** RT

Author Location RT (252-335)

Epitope Neutralizing yes Immunogen vaccine

Vector/Type: vaccinia Strain: B clade

HXB2 HIV component: RT

Species (Isotype) mouse

Ab Type RT thumb domain

References Ohba et al. 2001

5F: BALB/c mice were vaccinated with vaccinia carrying RT and a phage display library was produced and panned with RT – Fabs 5F and 5G were cloned, both recognizing an immunodominant neutralizing RT epitope in the region of the template primer-binding site in the thumb domain also recognized by MAb 7C4 – sequencing revealed the heavy chains and light chains of 7C4, 5G and 7C4 are related. Ohba *et al.* [2001]

No. 206

MAb ID 5G

**HXB2** Location RT

**Author Location** RT (252–335)

**Epitope** 

Neutralizing yes

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade HXB2 HIV component: RT

Species (Isotype) mouse

Ab Type RT thumb domain

References Ohba et al. 2001

5G: BALB/c mice were vaccinated with vaccinia carrying RT and a phage display library was produced and panned with RT – Fabs 5F and 5G were cloned, both recognizing an immunodominant neutralizing RT epitope in the region of the template primer-binding site in the thumb domain also recognized by MAb 7C4 – sequencing revealed the heavy chains and light chains of 7C4, 5G and 7C4 are related. Ohba *et al.* [2001]

**No.** 207

MAb ID 7C4

**HXB2 Location** RT

Author Location RT (252-335)

**Epitope** 

**Neutralizing** yes

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade

HXB2 HIV component: RT

Species (Isotype) mouse (IgG2a)

Ab Type RT thumb domain

References Ohba et al. 2001; Chiba et al. 1997; Chiba

et al. 1996

- 7C4: Fabs 5F and 5G both recognize the same immunodominant neutralizing RT epitope in the region of the template primer-binding site in the thumb domain recognized by MAb 7C4 sequencing revealed the heavy chains and light chains of 7C4, 5G and 7C4 are related. Ohba *et al.* [2001]
- 7C4: 7C4 inhibits RT from HIV-1 strains IIIB, Bru, and IMS-1 but not HIV-2 strains GH-1 and LAV-2, SIV MAC, nor SIV MND. Chiba et al. [1997]
- 7C4: 7C4 was produced from a hybridoma cell line derived from a BALB/c mouse repeatedly immunized with RT in a vaccinia construct, and was found to inhibit RT through binding to the template primer-binding site, a possible target for RT inhibitors. Chiba *et al.* [1996]

**HIV Antibodies Tables Integrase Antibodies** 

## **IV-C-9** Integrase Antibodies

No. 208

MAb ID 1C4

HXB2 Location Integrase (1-16)

**Author Location** Integrase (1–16 HXB2)

Epitope FLDGIDKAQDEHEKYH

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type N-term

References Nilsen et al. 1996; Haugan et al. 1995

- 1C4: One of a large set of MAbs that interact with the Nterminal part of integrase: 1C4, 2C11, 2E3, 3E11, 3F9, 5F8, 6G5, 7B6, 7C6 - these MAbs inhibit end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]
- 1C4: MAb interferes with integrase binding to DNA. Haugan et al. [1995]

No. 209

MAb ID 2C11

HXB2 Location Integrase (1–16)

**Author Location** Integrase (1–16 HXB2)

Epitope FLDGIDKAQDEHEKYH

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type N-term

References Nilsen et al. 1996

• 2C11: One of a large set of MAbs that interact with the N- Species (Isotype) mouse (IgG1 $\kappa$ ) terminal part of integrase: 1C4, 2C11, 2E3, 3E11, 3F9, 5F8, 6G5, 7B6, 7C6 - these MAbs inhibit end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]

**No.** 210

MAb ID 2E3

HXB2 Location Integrase (1-16)

**Author Location** Integrase (1–16 HXB2)

Epitope FLDGIDKAQDEHEKYH

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type N-term

References Ovod et al. 1992; Nilsen et al. 1996

• 2E3: One of a large set of MAbs that interact with the Nterminal part of integrase: 1C4, 2C11, 2E3, 3E11, 3F9, 5F8, 6G5, 7B6, 7C6 - these MAbs inhibit end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]

• 2E3: There are two MAbs called 2E3 – the other one binds to Nef. Ovod et al. [1992]

No. 211

MAb ID 3E11

HXB2 Location Integrase (1–16)

**Author Location** Integrase (1–16 HXB2)

Epitope FLDGIDKAQDEHEKYH

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

**Ab Type** N-term

References Nilsen et al. 1996; Otteken et al. 1992

- 3E11: One of a large set of MAbs that interact with the Nterminal part of integrase: 1C4, 2C11, 2E3, 3E11, 3F9, 5F8, 6G5, 7B6, 7C6 - these MAbs inhibit end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]
- 3E11: There is another MAb with this ID that recognizes p17. Otteken et al. [1992]
- 3E11: Recognized an epitope present on HIV-2/SIVmac, SIVagm, HIV-1, and SIVmnd. Otteken et al. [1992]

No. 212

MAb ID 3F9

HXB2 Location Integrase (1–16)

**Author Location** Integrase (1–16 HXB2)

Epitope FLDGIDKAQDEHEKYH

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

Ab Type N-term

References Nilsen et al. 1996

• 3F9: One of a large set of MAbs that interact with the Nterminal part of integrase: 1C4, 2C11, 2E3, 3E11, 3F9, 5F8, 6G5, 7B6, 7C6 - these MAbs inhibit end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]

**No.** 213

MAb ID 5F8

HXB2 Location Integrase (1-16)

**Author Location** Integrase (1–16 HXB2)

Epitope FLDGIDKAQDEHEKYH

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type N-term

References Nilsen et al. 1996; Haugan et al. 1995

• 5F8: There is another MAb with this ID that recognizes and unknown protein in HIV.

**HIV Antibodies Tables Integrase Antibodies** 

terminal part of integrase: 1C4, 2C11, 2E3, 3E11, 3F9, 5F8, 6G5, 7B6, 7C6 - these MAbs inhibit end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]

• 5F8: MAb interferes with integrase binding to DNA. Haugan et al. [1995]

**No.** 214

MAb ID 6G5

HXB2 Location Integrase (1–16)

**Author Location** Integrase (1–16 HXB2)

Epitope FLDGIDKAQDEHEKYH

Subtype B

Neutralizing no

Immunogen vaccine

*Vector/Type*: protein *Strain*: B clade HXB2 **Species (Isotype)** mouse (IgG1κ)

HIV component: Int

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type N-term

References Nilsen et al. 1996

• 6G5: One of a large set of MAbs that interact with the Nterminal part of integrase: 1C4, 2C11, 2E3, 3E11, 3F9, 5F8, 6G5, 7B6, 7C6 - these MAbs inhibit end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]

No. 215

MAb ID 7B6

**HXB2 Location** Integrase (1–16)

Author Location Integrase (1-16 HXB2)

Epitope FLDGIDKAQDEHEKYH

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type N-term

References Nilsen et al. 1996

• 7B6: One of a large set of MAbs that interact with the Nterminal part of integrase: 1C4, 2C11, 2E3, 3E11, 3F9, 5F8, 6G5, 7B6, 7C6 - these MAbs inhibit end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]

**No.** 216

MAb ID 7C6

HXB2 Location Integrase (1-16)

Author Location Integrase (1–16 HXB2)

Epitope FLDGIDKAQDEHEKYH

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type N-term

References Nilsen et al. 1996

• 5F8: One of a large set of MAbs that interact with the N- • 7C6: One of a large set of MAbs that interact with the Nterminal part of integrase: 1C4, 2C11, 2E3, 3E11, 3F9, 5F8, 6G5, 7B6, 7C6 – these MAbs inhibit end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]

No. 217

MAb ID 6C5

HXB2 Location Integrase (17-38)

Author Location Integrase (17-38 HXB2)

Epitope SNWRAMASDFNLPPVVAKEIVA

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

Ab Type N-term

References Nilsen et al. 1996; Haugan et al. 1995

- 6C5: This MAb inhibits end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]
- 6C5: MAb interferes with integrase binding to DNA. Haugan et al. [1995]

No. 218

MAb ID 8G4

HXB2 Location Integrase (22–31)

**Author Location** Integrase (12–42 HXB2)

Epitope MASDFNLPPV+GYIEAEVIPAETGQETAYFI?

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Nilsen et al. 1996; Haugan et al. 1995

- 8G4: This MAb reacted strongly with peptides IN(12-31) and IN(22-42), and less strongly with peptide IN(82-101) – it did not react with a deletion mutant of positions 17-38 - this MAb inhibits end processing and DNA joining, but had little effect on integration activities. Nilsen et al. [1996]
- 8G4: MAb interferes with integrase binding to DNA. Haugan et al. [1995]

No. 219

**MAb ID** 17 (mAb17)

HXB2 Location Integrase (25-35)

**Author Location** Integrase (25–35)

**Epitope** DFNLPPVVAKE

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG1)

References Yi et al. 2000; Levy-Mintz et al. 1996; Bizub-

Bender et al. 1994

• 17: Epitope mapped to helix-turn-helix motif in the N-term domain of Integrase, positions 25-35 - Zn binding stabilizes the Integrase-mAb17 complex - both MAb and Fab form of mAb17 inhibit Integrase activity - epitope region likely to be involved in protein-protein interaction. Yi et al. [2000]

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• 17: Used for the creation of single chain variable antibody fragments (SFvs) for internal cellular expression – neutralization of IN activity prior to integration, whether the Ab is expressed in the nucleolus or the cytoplasm – relative binding affinity to IN: 12 > 17 = 33 > 21 > 4. Levy-Mintz *et al.* [1996]

17: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized – the Zn finger motif is in the binding region – MAbs 14 and 17 form a competition group. Bizub-Bender *et al.* [1994]

No. 220

MAb ID 4D6

**HXB2 Location** Integrase (42–55)

**Author Location** Integrase (42–55 HXB2)

Epitope KCQLKGEAMHGQVD

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type N-term

References Nilsen et al. 1996; Haugan et al. 1995

- 4D6: This MAb inhibits end processing and DNA joining, and reduces reintegration activity. Nilsen et al. [1996]
- 4D6: MAb interferes with integrase binding to DNA. Haugan *et al.* [1995]

No. 221

**MAb ID** 7-16 (7-19)

**HXB2 Location** Integrase (50–159)

**Author Location** Integrase (50–159 HXB2)

**Epitope** 

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: chimeric maltose binding protein (MBP) Strain: B clade IIIB HIV com-

ponent: Int

Species (Isotype) mouse (IgG2b)

Ab Type Integrase catalytic core

Research Contact Yoshihiro Kitamura, Div of Mol Genetics,

Nat Inst of Infectious Diseases, Musashimu-

rayama, Japan

References Ishikawa et al. 1999

• 7-16: Binds to the central catalytic domain – the paper seems to sometimes call this antibody 7-16, sometimes 7-19, a possible typo. Ishikawa *et al.* [1999]

No. 222

MAb ID 4F6

HXB2 Location Integrase (56–102)

Author Location Integrase (56–102 HXB2)

Epitope CSPGIWQLDCTHLEGKVILVAVHVASGYIEAE-

VIPAETGQETAYFLL

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type Integrase catalytic core

References Nilsen et al. 1996; Haugan et al. 1995

- 4F6: MAb binding had minimal effects on IN in vitro activities.
   Nilsen et al. [1996]
- 4F6: MAb interferes with integrase binding to DNA. Haugan et al. [1995]

No. 223

MAb ID anti-K159

HXB2 Location Integrase (151–163)

**Author Location** Integrase (163–175)

Epitope VESMNKELKKIIG

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide HIV component: Int

Species (Isotype) rabbit (IgG)

References Maksiutov et al. 2002; Maroun et al. 1999

- anti-K159: This epitope is similar to a fragment of the human protein Apoptosis regulator BCL-W (KIAA0271), ESVNKE-MEPLVGQV. Maksiutov et al. [2002]
- anti-K159: Both the peptide K159, SQGVVESMNKELKKI-IGQVRDQAEHLKTA, and the Abs raised against this peptide inhibit Integrase activity K159 was found to fulfill condition of minimal number of helical heptads to achieve the formation of a stable coiled-coil structure Integrase is proposed to function as a dimer interacting in this region. Maroun *et al.* [1999]

**No.** 224

MAb ID 5D9

HXB2 Location Integrase (186–250)

**Author Location** Integrase (186–250 HXB2)

Epitope

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type Integrase DNA binding domain

References Nilsen et al. 1996; Haugan et al. 1995

- 5D9: MAb binding had minimal effects on IN *in vitro* activities. Nilsen *et al.* [1996]
- 5D9: While C-term and N-term anti-Integrase MAbs interfere with Integrase-DNA binding, 5D9 which binds more centrally, does not. Haugan *et al.* [1995]

No. 225

**MAb ID** 8-6

HXB2 Location Integrase (211–227)

**Author Location** Integrase (211–227 HXB2)

Epitope KELQKQITKIQNFRVYY

Subtype B

Neutralizing no

Immunogen vaccine

**HIV Antibodies Tables Integrase Antibodies** 

> Vector/Type: chimeric maltose binding protein (MBP) Strain: B clade IIIB HIV component: Int

Species (Isotype) mouse (IgG1)

Research Contact Yoshihiro Kitamura, Div of Mol Genetics, Nat Inst of Infectious Diseases, Musashimurayama, Japan

References Ishikawa et al. 1999

• 8-6: Antibody binds proximal to the DNA binding region. Ishikawa *et al*. [1999]

**No.** 226

**MAb ID** 19 (2-19, scAb2-19)

HXB2 Location Integrase (228–236) **Author Location** Integrase (228–236 LAI)

Epitope RDSRNPLWK

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG1)

References Kitamura et al. 1999; Levy-Mintz et al. 1996; Bizub-Bender et al. 1994

- 19: Called 2-19, scAb2-19 is a single-chain Ab made from MAb 2-19 -acts intra-cellularly to block infection at low MOI by binding to integrase - scAb interfered with the folding of Gag-Pol polyprotein, the Ab did not affect viral production in LAI transfected cells, but the virus produced was less infectious – authors suggest that the epitope may be conformational. Kitamura et al. [1999]
- 19: BALBc mice were immunized with rec integrase, hybrido- Research Contact Yoshihiro Kitamura, Div of Mol Genetics, mas expressing anti-integrase Abs were generated, and the antibodies characterized – 19 has a low binding affinity. Bizub-Bender et al. [1994]

No. 227

**MAb ID** 2-19

HXB2 Location Integrase (228–236)

**Author Location** Integrase (228–236 HXB2)

**Epitope** RDSRNPLWK

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: chimeric maltose binding protein (MBP) Strain: B clade IIIB HIV component: Int

Species (Isotype) mouse (IgG2b)

Ab Type Integrase DNA binding domain

Research Contact Yoshihiro Kitamura, Div of Mol Genetics, Nat Inst of Infectious Diseases, Musashimu-

rayama, Japan

References Ishikawa et al. 1999

• 2-19: MAb inhibits RT-Integrase interaction, and the terminal cleavage and strand transfer functions of Integrase, but not the disintegration activity. Ishikawa et al. [1999]

No. 228

**MAb ID** 8-22

HXB2 Location Integrase (237–252)

Author Location Integrase (237–252 HXB2)

Epitope GPAKLLWKGEGAVVIQ

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: chimeric maltose binding protein (MBP) Strain: B clade IIIB HIV com-

ponent: Int

Species (Isotype) mouse (IgG1)

Ab Type Integrase DNA binding domain

Research Contact Yoshihiro Kitamura, Div of Mol Genetics,

Nat Inst of Infectious Diseases, Musashimu-

rayama, Japan

References Ishikawa et al. 1999

• 8-22: MAb inhibits the terminal cleavage and strand transfer functions of Integrase, but not the disintegration activity. Ishikawa et al. [1999]

No. 229

**MAb ID** 4-20

HXB2 Location Integrase (253–261)

**Author Location** Integrase (253–261 HXB2)

Epitope DNSDIKVVP

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: chimeric maltose binding protein (MBP) Strain: B clade IIIB HIV com-

ponent: Int

Species (Isotype) mouse (IgG1)

Ab Type Integrase DNA binding domain

Nat Inst of Infectious Diseases, Musashimu-

rayama, Japan

References Ishikawa et al. 1999

• 4-20: Inhibits the terminal cleavage and strand transfer functions of Integrase, but not the disintegration activity. Ishikawa et al. [1999]

No. 230

**MAb ID** 6-19

HXB2 Location Integrase (262-270)

**Author Location** Integrase (261–270 HXB2)

Epitope RRKAKIIRD

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: chimeric maltose binding protein (MBP) Strain: B clade IIIB HIV com-

ponent: Int

Species (Isotype) mouse (IgG2b)

Ab Type Integrase DNA binding domain

Research Contact Yoshihiro Kitamura, Div of Mol Genetics, Nat Inst of Infectious Diseases, Musashimu-

rayama, Japan

References Ishikawa et al. 1999

• 6-19: Inhibits the terminal cleavage and strand transfer functions of Integrase, but not the disintegration activity. Ishikawa et al. [1999]

No. 231

Pol Antibodies HIV Antibodies Tables

MAb ID 7C3

**HXB2 Location** Integrase (262–271)

**Author Location** Integrase (262–271 HXB2)

Epitope RRKAKIIRDY

Subtype B Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Nilsen et al. 1996; Haugan et al. 1995

- 7C3: A set of three MAbs recognize an epitope in this region, 7C3, 7F11, and 8E5 all three HIV-1 MAbs cross-react with HIV-2 IN these MAbs inhibit end-processing, DNA joining and reintegration, and had little effect on disintegration. Nilsen *et al.* [1996]
- 7C3: MAb interferes with integrase binding to DNA. Haugan *et al.* [1995]

No. 232

MAb ID 7F11

HXB2 Location Integrase (262–271)

Author Location Integrase (262–271 HXB2)

Epitope RRKAKIIRDY

**Subtype** B **Neutralizing** no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Lasky et al. 1987; Nilsen et al. 1996

- 7F11: A set of three MAbs recognize an epitope in this region, 7C3, 7F11, and 8E5 all three HIV-1 MAbs cross-react with HIV-2 IN these MAbs inhibit end-processing, DNA joining and reintegration, and had little effect on disintegration. Nilsen *et al.* [1996]
- 7F11: There is another MAb with this name that binds to gp120. Lasky *et al.* [1987]

**No.** 233

MAb ID 8E5

**HXB2 Location** Integrase (262–271)

**Author Location** Integrase (262–271 HXB2)

Epitope RRKAKIIRDY

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade HXB2

HIV component: Int

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Nilsen et al. 1996; Haugan et al. 1995

- 8E5: A set of three MAbs recognize an epitope in this region, 7C3, 7F11, and 8E5 all three HIV-1 MAbs cross-react with HIV-2 IN these MAbs inhibit end-processing, DNA joining and reintegration, and had little effect on disintegration. Nilsen *et al.* [1996]
- 8E5: MAb interferes with integrase binding to DNA. Haugan et al. [1995]

**No.** 234

MAb ID MAb 35

HXB2 Location Integrase (264–273)

**Author Location** Integrase (264–273)

Epitope KAKIIRDYGK

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

**Species** (**Isotype**) mouse ( $IgG\kappa$ )

References Acel et al. 1998; Barsov et al. 1996

- MAb 35: Integrase was shown to have intrinsic DNA polymerase activity that can catalyze gap repair MAb 35 inhibits this activity. Acel *et al.* [1998]
- MAb 35: There appears to be two different IN Abs with similar names: MAb 35 and 35. Barsov et al. [1996]
- MAb 35: Although MAb 35 does not inhibit HIV-1 IN, Fab 35 inhibits 3'-end processing, strand transfer and disintegration. Barsov et al. [1996]

No. 235

MAb ID polyclonal

**HXB2 Location** Integrase

**Author Location** Integrase

**Epitope** 

Subtype multiple

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Barin et al. 2005

Country France

Keywords acute/early infection, assay development

• A combination of 4 antigenic regions was used to differentiate between early (<180 days) and chronic infection. These regions were: p24; the gp41 peptide spanning the immunodominant epitope (IDE) of gp41, RVAVERYLKDQQLLGIWGC-SGKICTTAV, and a subtype D version of this peptide; 5 V3 consensus peptides including A, B, C, D, and CRF01-AE; and Integrase. V3 and the IDE provide the best discrimination, with >20 fold higher levels in chronic infection when assayed by EIA using dried serum spots. Antibodies to Integrase and p24 were not as distinctive, and people tend to lose, not increase, responses to p24 over time. Integrase antibodies are among the last to appear after infection. This assay can be used to identify samples from early infection with high sensitivity and specificity. Barin *et al.* [2005] (assay development, acute/early infection)

# **IV-C-10** Pol Antibodies

No. 236

MAb ID 12

HXB2 Location Pol

**Author Location** Integrase (1–58)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG2a)

HIV Antibodies Tables Pol Antibodies

**References** Levy-Mintz *et al.* 1996; Bizub-Bender *et al.* 1994

• 12: Used for the creation of single-chain variable antibody fragments (SFvs) for internal cellular expression – neutralization of IN activity prior to integration, whether the Ab is expressed in the nucleolus or the cytoplasm – relative binding affinity to IN: 12 > 17 = 33 > 21 > 4. Levy-Mintz *et al.* [1996]

• 12: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized – the Zn finger motif is in the binding region – MAbs 12, 13 and 35 form a competition group. Bizub-Bender *et al.* [1994]

No. 237

**MAb ID** 13

HXB2 Location Pol

Author Location Integrase (1-58)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG1)

References Bizub-Bender et al. 1994

13: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized – the Zn finger motif is in the binding region – MAbs 12, 13 and 35 form a competition group. Bizub-Bender et al. [1994]

**No.** 238

MAb ID 14

**HXB2 Location** Pol

**Author Location** Integrase (1–58)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

**Species (Isotype)** mouse (IgG1)

References Bizub-Bender et al. 1994

• 14: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized – the Zn finger motif is in the binding region – MAbs 14 and 17 form a competition group. Bizub-Bender *et al.* [1994]

No. 239

**MAb ID** 16

HXB2 Location Pol

**Author Location** Integrase

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG2a)

References Bizub-Bender et al. 1994

16: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized. Bizub-Bender et al. [1994]

**No.** 240

MAb ID 1C12B1

HXB2 Location Pol

**Author Location** RT (431–521)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: RT

Species (Isotype) mouse

References Ferns et al. 1991

- 1C12B1: UK Medical Research Council AIDS reagent: ARP384.
- 1C12B1: Recognized both p66 and p51 in Western blot, binds to C terminus. Ferns *et al.* [1991]

No. 241

MAb ID 21

HXB2 Location Pol

**Author Location** Integrase (58–141)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG2b)

**References** Levy-Mintz *et al.* 1996; Bizub-Bender *et al.* 1994

- 21: Used for the creation of single chain variable antibody fragments (SFvs) for internal cellular expression neutralization of IN activity prior to integration, whether the Ab is expressed in the nucleolus or the cytoplasm relative binding affinity to IN: 12 > 17 = 33 > 21 > 4. Levy-Mintz *et al.* [1996]
- 21: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized. Bizub-Bender *et al.* [1994]

No. 242

**MAb ID** 32 (mAb32, Fab32)

HXB2 Location Pol

Author Location Integrase (223–266)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG2b)

**References** Yi *et al.* 2002; Yi & Skalka 2000; Bizub-Bender *et al.* 1994

- 32: Called mAb32 mAb33 and mAb32 compete for binding to the C-term domain of Integrase – while mAb32 only weakly inhibits IN activity, mAb33 inhibits strongly, mAb32 has a lower affinity than mAb33, and Fab32 does not inhibit at all while Fab33 inhibits DNA binding a catalytic activity. Yi et al. [2002]
- 32: Limited proteolysis combined with mass spectrometric analysis indicates Fab32 binds to two strands of the beta sheet, beta1 223F, 224R, 226Y, and 228R and beta5 264K and 266K. Yi & Skalka [2000]
- 32: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized MAbs 32 and 33 form a competition group. Bizub-Bender *et al.* [1994]

Pol Antibodies **HIV Antibodies Tables** 

No. 243

MAb ID 35

**HXB2 Location** Pol

Author Location Integrase (1-58)

**Epitope** Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG2b)

References Bizub-Bender et al. 1994

- 35: There appears to be two IN Abs with similar names: MAb 35 and 35. Bizub-Bender et al. [1994]
- 35: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized - the Zn finger motif is in the binding region - MAbs 12, 13 and 35 form a competition group. Bizub-Bender et al. [1994]

No. 244

MAb ID 3D12

**HXB2 Location** Pol

**Author Location RT** 

**Epitope** 

Neutralizing Immunogen vaccine

Vector/Type: vaccinia HIV component: RT

Species (Isotype) mouse (IgG2a)

References Chiba et al. 1997

• 3D12: There is an anti-Nef MAb that also has this name (see Chiba et al. [1997]) Chiba et al. [1997]

No. 245

MAb ID 3F10

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

Neutralizing Immunogen vaccine

Vector/Type: vaccinia HIV component: RT

Species (Isotype) mouse (IgG2a)

References Chiba et al. 1997

No. 246

MAb ID 4

HXB2 Location Pol

Author Location Integrase (141-172)

**Epitope** Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG2b)

References Levy-Mintz et al. 1996; Bizub-Bender et al. Research Contact B. Ferns and R. Tedder 1994

- 4: Used for the creation of single chain variable antibody fragments (SFvs) for internal cellular expression – neutralization of IN activity prior to integration, whether the Ab is expressed in the nucleolus or the cytoplasm – relative binding affinity to IN: 12 > 17 = 33 > 21 > 4. Levy-Mintz *et al.* [1996]
- 4: There is another MAb with this ID that reacts with gp41. Bizub-Bender et al. [1994]

• 4: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized – 4 has a low binding affinity. Bizub-Bender et al. [1994]

No. 247

MAb ID 6B9

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia HIV component: RT

Species (Isotype) mouse (IgG2a)

References Chiba et al. 1997

**No.** 248

MAb ID 7C4

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia HIV component: RT

Species (Isotype) mouse (IgG1)

References Chiba et al. 1997

• 7C4: Dose-dependent inhibition of polymerase activity of RT of strains IIIB, Bru and IMS-1, but not HIV-2 strains GH-1 or LAV-2 or SIV strains MAC or MND. Chiba et al. [1997]

No. 249

MAb ID RT-4

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

Neutralizing no

**Immunogen** 

Species (Isotype) mouse (IgG2b)

References Gu et al. 1996; Li et al. 1993

• RT-4: Increased nevirapine and delavirdine inhibition, no effect on AZT inhibition. Gu et al. [1996]

No. 250

MAb ID RT70

**HXB2 Location** Pol

Author Location RT (231–315)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: RT

Species (Isotype) mouse (IgG1)

**References** Ferns *et al.* 1991

- RT7O: UK Medical Research Council AIDS reagent: ARP381.
- RT7O: Conformational epitope located centrally in the protein - inhibited RT enzyme activity and thus may bind close to the active site of the enzyme. Ferns et al. [1991]

No. 251 MAb ID RT7U HIV Antibodies Tables Pol Antibodies

HXB2 Location Pol

Author Location RT (231-315)

**Epitope Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: RT

Species (Isotype) mouse

Research Contact B. Ferns and R. Tedder

References Ferns et al. 1991

• RT7U: UK Medical Research Council AIDS reagent: ARP380.

• RT7U: Has a conformational epitope – reacts with p66 and p51 in WB. Ferns *et al.* [1991]

No. 252

MAb ID anti-HIV-1 RT

HXB2 Location Pol

**Author Location RT** 

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) mouse (IgG)

**References** Wainberg & Gu 1995; Maciejewski *et al.* 1995; di Marzo Veronese *et al.* 1986

- anti-HIV-1 RT: Cloned heavy and light chains to express Fab intracellularly, preventing HIV infection in vitro – this MAb was broadly cross-reactive with clinical strains and even HIV-2. Maciejewski et al. [1995]
- Commentary on Maciejewski et al. Wainberg & Gu [1995]

**No.** 253

MAb ID polyclonal

**HXB2 Location** Pol

**Author Location** p55

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: virus-like particle (VLP) HIV

component: Gag, gp120, V3

Species (Isotype) macaque

References Wagner et al. 1998b

A VLP is a non-infectious virus-like particle self-assembled from HIV Pr55 gag – macaques were immunized with VLPs bound to either gp120 or V3+CD4 linear domains – gag and env CTL specific CTL were stimulated in each case, and Ab response to gag and gp120 and was elicited, but the gp120 neutralizing response occurred only with whole gp120, not V3+CD4 – despite the CTL and Ab response, immunized macaques were infected by intervenous challenge with SHIV chimeric challenge stock. Wagner et al. [1998b]

No. 254

MAb ID polyclonal

HXB2 Location Pol

**Author Location RT** 

Epitope

Neutralizing

Immunogen vaccine

Vector/Type: DNA HIV component: Env,

Gag, Pol, Vif Adjuvant: B7, IL-12

Species (Isotype) mouse

References Kim et al. 1997b

 A gag/pol, vif or CMN160 DNA vaccine, when delivered in conjunction with the plasmid encoding the co-stimulatory molecules B7 and IL-12, gave a dramatic increase in both the cytotoxic and proliferative responses in mice, as well as Ab response detected by ELISA. Kim et al. [1997b]

**No.** 255

MAb ID polyclonal

HXB2 Location Pol

Author Location RT (203-219)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: Salmonella HIV component:

RT

Species (Isotype) mouse (IgA)

References Burnett et al. 2000

 A live attenuated bacterial vaccine, Salmonella SL3261pHART, with an inserted HIV RT gene fragment in the Lpp-OmpA-HIV fusion protein, induced a lymphoproliferative Th response and fecal RT-specific IgA in BALB/c mice. Burnett et al. [2000]

No. 256

MAb ID 33 (mAb33, Fab33, 33D5, mab 33)

HXB2 Location Pol

Author Location Integrase (223–268 HXB2)

**Epitope** 

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Int

Species (Isotype) mouse (IgG2b)

Ab Type C-term

References Schreiber et al. 2005; Yi et al. 2002; Yi &

Skalka 2000; Levy-Mintz et al. 1996; Bizub-

Bender et al. 1994

**Keywords** antibody binding site definition and exposure,

computational epitope prediction, mimotopes,

structure

• 33: Called Fab33. A new computer program designed to recognize conformational epitopes in 3D structures that correspond to linear peptide mimotopes (3DEX) was tested using the known conformational epitope of this Fab. 223F, 224R, 226Y, 244K, 267I, and 268I, previously defined as the epitope from NMR structural studies (Yi2002) were confirmed, along with two additional amino acids, (A265 and K266). Schreiber et al. [2005] (antibody binding site definition and exposure, mimotopes, computational epitope prediction, structure)

33: Called mAb33 – mAb33 and mAb32 compete for binding to the C-term domain of Integrase – while mAb32 only weakly inhibits IN activity, mAb33 inhibits strongly, mAb32 has a lower affinity than mAb33, and Fab32 does not inhibit at all while Fab33 inhibits catalytic activity and DNA binding – heteronuclear NMR indicated eight residues of Integrase are immobilized upon Fab33 binding, two in the core of the protein, and 6 on the outer face that form a contiguous patch likely to contain the epitope – 223F, 224R, 226Y, 244K, 267I, and 268I, which may be a useful target for drug design – the Fab33-IN

Vif Antibodies HIV Antibodies Tables

complex is far more soluble than IN alone and may be useful for crystallization. Yi *et al.* [2002] (antibody binding site definition and exposure)

- 33: Limited proteolysis combined with mass spectrometric analysis were used to define the binding site for Fab32, but Fab33 binding to the Intergrase C-term domain left it resistant to proteolytic digestion. Yi & Skalka [2000]
- 33: Used for the creation of single chain variable antibody fragments (SFvs) for internal cellular expression neutralization of IN activity prior to integration, whether the Ab is expressed in the nucleolus or the cytoplasm relative binding affinity to IN: 12 > 17 = 33 > 21 > 4. Levy-Mintz *et al.* [1996]
- 33: BALBc mice were immunized with rec integrase, hybridomas expressing anti-integrase Abs were generated, and the antibodies characterized MAbs 32 and 33 form a competition group. Bizub-Bender *et al.* [1994]

No. 257

MAb ID F-6

HXB2 Location Pol

Author Location RT (BH-10)

**Epitope** 

Subtype B

Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human

Ab Type C-term

Research Contact Amon Hizi, Sackler School of Medicine, Tel

Aviv Isreal

References Herschhorn et al. 2003

Keywords antibody generation, antibody sequence, vari-

able domain, immunotherapy

• F-6: One of five human single chain Fv (ScFv) Abs isolated from an phage display library. F-6 was shown to bind to the carboxyl terminal segment of the p66 RT polypeptide, corresponding to RNase H. F-6 inhibited the DNA and RNA-dependent DNA polymerase (RDDP) activities of HIV-1 RT; two others, 6E9 and 5B11, inhibited RDDP and so have possible therapeutic potential. To pinpoint the mechanism of inhibition, three peptides were synthesized corresponding to the CDR3 sequences of F-6, and a cyclic version of the CDR H3 region bound to purified RT and blocked RDDP. Herschhorn et al. [2003] (antibody generation, immunotherapy, antibody sequence, variable domain)

#### IV-C-11 Vif Antibodies

No. 258

MAb ID TG002

HXB2 Location Vif (34–47)

**Author Location** Vif (34–47)

Epitope KARGWFYRHHYESP?

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Vif

Species (Isotype) mouse Research Contact Transgene

References

 TG002: This MAb was raised in response to a rec Vif protein derived from E. coli.

 TG002: NIH AIDS Research and Reference Reagent Program: 2746

No. 259

MAb ID TG001

**HXB2 Location** Vif (176–192)

**Author Location** Vif (176–192)

Epitope KPQKTKGHRGSHTMNGH?

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Vif

Species (Isotype) mouse

Ab Type C-term

Research Contact Transgene

#### References

- TG001: This antibody was raised in response to a rec Vif protein derived from E. coli.
- TG001: NIH AIDS Research and Reference Reagent Program: 2745.

No. 260

MAb ID J4

**HXB2 Location** Vif

**Author Location (HXB2)** 

**Epitope** 

Subtype B

**Neutralizing** 

Immunogen

Species (Isotype) humanized rabbit

References Goncalves et al. 2002

• J4: The authors developed a Vif-specific intrabody singlechain FAb fragment of J4 called 14BL. When expressed intracellularly in the cytoplasm this intrabody efficiently bound Vif protein and neutralized its infectivity enhancing function. Intrabody-expressing transduced cells were highly refractory to challenge with the laboratory strain NL43 and with primary isolate strains of HIV-1. Goncalves *et al.* [2002]

**No.** 261

MAb ID polyclonal

**HXB2 Location** Vif

**Author Location** Vif

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: DNA HIV component: Env, Gag, Pol, Vif Adjuvant: B7, IL-12

Species (Isotype) mouse

**References** Kim *et al.* 1997b

 A gag/pol, vif or CMN160 DNA vaccine, when delivered in conjunction with the plasmid encoding the co-stimulatory molecules B7 and IL-12, gave a dramatic increase in both the cytotoxic and proliferative responses in mice, as well as an Ab response detected by ELISA. Kim et al. [1997b]

## **IV-C-12** Vpr Antibodies

No. 262

MAb ID polyclonal

**HXB2 Location** Vpr

Author Location Vpr (89.6)

Epitope Subtype B Neutralizing

Immunogen HIV-1 infection Species (Isotype) human (IgG)

References Richardson et al. 2003

**Country** France

Keywords rate of progression

• Serum samples were obtained from the French GRIV (genetic resistance to HIV) cohort and tested for anti-Tat and anti-Vpr responses, as both may contribute as extracellular proteins to pathogenesis. Serum anti-Vpr IgG responses were significantly higher in stable non-progressors (CD4+ T cell counts greater than 500 cell/ul after being positive for 8 years with no ART) and unstable non-progressors (declined during a 20 month follow up), than fast progressors (CD4+ T less than 300 cells/ul within 2 years of seroconversion, some HAART). Serum anti-Tat IgG was found to be significantly higher in stable non-progressors compared to unstable non-progressors and fast progressors indicating that higher levels of serum anti-Tat IgG are associated with maintenance of non-progression status. Richardson *et al.* [2003] (rate of progression)

### **IV-C-13** Tat Antibodies

No. 263

MAb ID polyclonal

HXB2 Location Tat (1–15)

**Author Location** Tat (1–15 89.6)

Epitope MEPVDRPLEPWKHPG

Subtype B Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6, B clade HXBc2 HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (Isotype) macaque (IgG)

Ab Type C-term, N-term, Tat basic region

References Silvera et al. 2002

**Keywords** antibody binding site definition and exposure, vaccine antigen design

• Anti-Tat and Tat toxoid responses were raised in rhesus macaques using HxBc2 and 89.6P Tat, and Tat toxoids.High anti-Tat IgG responses were observed in all animals, and helper responses were detected in 8/16, and IFN gamma CTL in 11/16.. Ab and proliferative responses were observed, and the trucated 86 amino acid IIIB Tat was more immunogenic than the full 102 amino acid Tat. Tat vaccinated animals when challenged with SHIV89.6 were infected despite a robust immune response. The most consistent reactions to the vaccinations were to peptides in regions: N-term, 1-15 (MEPVDRPLEP-WKHPG), basic domain 46-60 (SYGRKKRRQRRAHQ),

and 61-91, particularly C-term 76-90 (QPRGDPTGPKQKKK). Silvera *et al.* [2002] (**antibody binding site definition and exposure**, **vaccine antigen design**)

No. 264

MAb ID polyclonal

HXB2 Location Tat (1-20)

**Author Location** Tat (1–20 IIIB BH10)

Epitope MEPVDPRLEPWKHPGSQPKT?

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA), macrophage activations in the Component of the Property of the Prop

ing lipopeptide-2 (MALP-2)

Species (Isotype) mouse (IgA, IgG)

References Borsutzky et al. 2003

**Keywords** adjuvant comparison, genital and mucosal immunity, immunodominance, mucosal immunity, Th1, Th2, vaccine-induced epitopes

• Intranasal immunization of BALB/c mice Tat with MALP-2 induced stronger immune responses than i.p. vaccination of Tat with IFA. Also i.n. MALP-2 vaccination favored Th1 responses, while i.p IFA favored Th2. Ab, T help and CTL responses were observed. MALP-2 enhanced Tat-specific IgA responses in the lung and in the genital tract. Three linear Ab epitopes were recognized. The most frequent response (80% of vaginal lavage, 100% of sera and lung lavage) was to Tat peptide 1-20. The second strongest to Tat 46-60 (20% of vaginal lavage, 50% of lung lavage, 50% of sera). Finally, 50% of the sera of Tat+ IFA i.p. immunized mice recognized Tat 73-86. Borsutzky et al. [2003] (adjuvant comparison, genital and mucosal immunity, vaccine-induced epitopes, immunodominance, mucosal immunity, Th1, Th2)

No. 265

MAb ID TA9

HXB2 Location Tat (1-20)

Author Location Tat (1-20 Lai/Bru)

Epitope MEPVDPRLEPGSQPKT

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU HIV component: Tat Adjuvant: Complete

Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgG)

**Ab Type** N-term

Research Contact Dr. J.-L. Guesdon, Institut Pasteur, Paris

References Belliard et al. 2003

**Keywords** subtype comparisons

• This is one of 5 anti-Tat murine monoclonal antibodies generated in this study. TA9 is clade B specific, and does not recognize Tat from clade A, C, D, or CRF01 (AE).TA9 binds to the Tat peptide aa 1-61 strongly, and is also able to bind to Tat aa 1-20, and Tat peptide aa 8-53. Belliard *et al.* [2003] (subtype comparisons)

**No.** 266

Tat Antibodies HIV Antibodies Tables

MAb ID TD84
HXB2 Location Tat (1–20)
Author Location Tat (1–20 Lai/Bru)
Epitope MEPVDPRLEPGSQPKT
Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU HIV component: Tat Adjuvant: Complete Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgG)

Ab Type N-term

Research Contact Dr. J.-L. Guesdon, Institut Pasteur, Paris

**References** Belliard *et al.* 2003 **Keywords** subtype comparisons

• This is one of 5 anti-Tat murine monoclonal antibodies generated in this study. TD84 is clade B specific, and does not recognize Tat from clade A, C, D, or CRF01 (AE). It reacts strongly with aa 1-61, and is able to react with aa 1-20. Belliard *et al.* [2003] (**subtype comparisons**)

No. 267
MAb ID TE135
HXB2 Location Tat (1–20)
Author Location Tat (1–20 Lai/Bru)
Epitope MEPVDPRLEPGSQPKT
Subtype B
Neutralizing

Vector/Type: protein Strain: B clade BRU HIV component: Tat Adjuvant: Complete Freund's Adjuvant (CFA)

 $\boldsymbol{Species}\;(\boldsymbol{Isotype})\;\;\text{mouse}\;(\boldsymbol{IgG})$ 

Immunogen vaccine

Ab Type N-term

Research Contact Dr. J.-L. Guesdon, Institut Pasteur, Paris

**References** Belliard *et al.* 2003 **Keywords** subtype comparisons

• This is one of 5 anti-Tat murine monoclonal antibodies generated in this study. TE135 is clade B specific, and does not recognize Tat from clade A, C, D, or CRF01 (AE). It reacts strongly with aa 1-61, and is able to react with aa 1-20. Belliard *et al.* [2003] (**subtype comparisons**)

**No.** 268

**MAb ID** polyclonal **HXB2 Location** Tat (1–24)

**Author Location** Tat (1–24)

Epitope MEPVDPRLEPWKHPGSQPKTACTN

Neutralizing

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade HIV component: Tat Adjuvant: Montanide (ISA

51)

Species (Isotype) human (IgG)

Ab Type N-term

References Noonan et al. 2003

**Keywords** immunotherapy, vaccine-specific epitope characteristics

• Intramuscular injection of Tat-toxoid induced high titers of anti-Tat reactivity in serum samples of six HIV-1 positive and of four HIV negative study subjects. Anti-Tat antibodies successfully blocked extracellular Tat from transactivating HIV Tat-sensitive promotors. The anti-Tat IgG response in sera from two healthy and HIV infected patients inhibited cell entry of synthetic Tat, thus blocking its functional activity. Additionally, the anti-Tat antibodies inhibited intercellular Tat transfer as demonstrated by a co-culture cell system. All HIV-1 infected patients had Ab responses to the N-term region of Tat, and 4/4 HIV-1 + and 5/6 HIV-1 negative patients responded to the basic domain. Several additional peptides were recognized either exclusively or more commonly in the HIV+ people. The N-terminus region of Tat mediates binding to CD26, that may be involved in modulation of chemokine function, and may also mediate Tcell apotosis. Noonan et al. [2003] (vaccine-specific epitope characteristics, immunotherapy)

**No.** 269

MAb ID NT3/2D1.1

HXB2 Location Tat (2-15)

**Author Location** Tat

Epitope EPVDPNLEPWNHPS

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: Tat

Species (Isotype) mouse (IgG1a)

Ab Type N-term

References Dingwall et al. 1989

- NT3/2D1.1: UK Medical Research Council AIDS reagent: ARP352.
- NT3/2D1.1: Immunoprecipitates and immunoblots HIV-1 tat protein. Dingwall *et al.* [1989]

No. 270

**MAb ID** 1.2

HXB2 Location Tat (2–17)

**Author Location** Tat (1–16)

Epitope EPVDPRLEWKHPGSQ

Neutralizing

**Immunogen** 

Species (Isotype)

References Ranki et al. 1995; Ovod et al. 1992

• 1.2: Weak expression of Tat observed in HIV+ brain tissue sample, in contrast to Nef. Ranki *et al.* [1995]

**No.** 271

**MAb ID** 1D9D5

HXB2 Location Tat (2-21)

**Author Location** Tat

Epitope EPVDPRLEWKHPGSQPKTA

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Tat

Species (Isotype) mouse (IgG1)

Ab Type N-term

References Valvatne et al. 1996; Mhashilkar et al. 1995

HIV Antibodies Tables Tat Antibodies

- 1D9D5: Exogenously delivered Tat can efficiently transactivate an HIV-LTR-CAT construct in HeLa cells in the presence of 1D9D5, suggesting when considered with the results of Mhashilkar *et al.* [1995], that free Tat and not Ab bound is taken up by cells Valvatne *et al.* [1996]. Mhashilkar *et al.* [1995]; Valvatne *et al.* [1996]
- 1D9D5: Single chain antibodies, intrabodies, were engineered that can be stably expressed in the cytoplasm of mammalian cells co-expression of an N-term intrabody can inhibit transactivation of an HIV LTR-CAT construct and block import into nucleus, but intrabody specific for exon 2 did not inhibit activity. Mhashilkar et al. [1995]

**No.** 272 **MAb ID** TB12

HXB2 Location Tat (44–60)

Author Location Tat (44-61 Lai/Bru)

Epitope GISYGRKKRRQRRPPQG

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU HIV component: Tat Adjuvant: Complete Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgG)

Ab Type Tat basic region

Research Contact Dr. J.-L. Guesdon, Institut Pasteur, Paris

**References** Belliard *et al.* 2003 **Keywords** subtype comparisons

• This is one of 5 anti-Tat murine monoclonal antibodies generated in this study. TB12 is clade B and D specific, and does not recognize Tat from clade A, C, or CRF01 (AE). It reacts strongly with aa 1-61, and is also able to react with aa 44-61, in the basic region involved in Tat uptake. Belliard *et al.* [2003] (subtype comparisons)

No. 273

MAb ID polyclonal

HXB2 Location Tat (46-60)

**Author Location** Tat (46–60 IIIB BH10)

Epitope SYGRKKRRQRRRAHQ?

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA), macrophage activating lipopeptide-2 (MALP-2)

Species (Isotype) mouse (IgA, IgG)

References Borsutzky et al. 2003

**Keywords** adjuvant comparison, genital and mucosal immunity, immunodominance, mucosal immunity, Th1, Th2, vaccine-induced epitopes

Intranasal immunization of BALB/c mice Tat with MALP-2 induced stronger immune responses than i.p. vaccination of Tat with IFA. Also i.n. MALP-2 vaccination favored Th1 responses, while i.p IFA favored Th2. Ab, T help and CTL responses were observed. MALP-2 enhanced Tat-specific IgA responses in the lung and in the genital tract. Three linear Ab epitopes were recognized. The most frequent response

(80% of vaginal lavage, 100% of sera and lung lavage) was to Tat peptide 1-20. The second strongest to Tat 46-60 (20% of vaginal lavage, 50% of lung lavage, 50% of sera). Finally, 50% of the sera of Tat+ IFA i.p. immunized mice recognized Tat 73-86. Borsutzky *et al.* [2003] (adjuvant comparison, genital and mucosal immunity, vaccine-induced epitopes, immunodominance, mucosal immunity, Th1, Th2)

No. 274

MAb ID polyclonal

HXB2 Location Tat (46–60)

**Author Location** Tat (46–60 89.6)

Epitope SYGRKKRRQRRRAHQ

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6, B clade HXBc2 HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (Isotype) macaque (IgG)

Ab Type C-term, N-term, Tat basic region

References Silvera et al. 2002

**Keywords** antibody binding site definition and exposure, vaccine antigen design

• Anti-Tat and Tat toxoid responses were raised in rhesus macaques using HxBc2 and 89.6P Tat, and Tat toxoids. High anti-Tat IgG responses were observed in all animals, and helper responses were detected in 8/16, and IFN gamma CTL in 11/16. Ab and proliferative responses were observed, and the trucated 86 amino acid IIIB Tat was more immunogenic than the full 102 amino acid Tat. Tat vaccinated animals when challenged with SHIV89.6 were infected despite a robust immune response. The most consistent reactions to the vaccinations were to peptides in regions: N-term, 1-15 (MEPVDRPLEPWKHPG), basic domain 46-60 (SYGRKKRQRRAHQ), and 61-91, particularly C-term 76-90 (QPRGDPTGPKQKKK). Silvera et al. [2002] (antibody binding site definition and exposure, vaccine antigen design)

No. 275

MAb ID polyclonal

HXB2 Location Tat (46–60)

**Author Location** Tat (46–60 89.6)

Epitope SYGRKKRRQRRRAHQ

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6, B clade HXBc2 HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (Isotype) macaque (IgG)

Ab Type C-term, N-term, Tat basic region

References Silvera et al. 2002

**Keywords** antibody binding site definition and exposure, vaccine antigen design

 Anti-Tat and Tat toxoid responses were raised in rhesus macaques using HxBc2 and 89.6P Tat, and Tat toxoids. High anti-Tat IgG responses were observed in all animals, and helper responses were detected in 8/16, and IFN gamma CTL in 11/16... Tat Antibodies HIV Antibodies Tables

Ab and proliferative responses were observed, and the trucated 86 amino acid IIIB Tat was more immunogenic than the full 102 amino acid Tat. Tat vaccinated animals when challenged with SHIV89.6 were infected despite a robust immune response. The most consistent reactions to the vaccinations were to peptides in regions: N-term, 1-15 (MEPVDRPLEP-WKHPG), basic domain 46-60 (SYGRKKRQRRAHQ), and 61-91, particularly C-term 76-90 (QPRGDPTGPKQKKK). Silvera *et al.* [2002] (antibody binding site definition and exposure, vaccine antigen design)

No. 276
MAb ID polyclonal

**HXB2 Location** Tat (47–60) **Author Location** Tat (46–60)

Epitope YGRKKRRQRRRPP0

**Neutralizing** 

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade HIV component: Tat Adjuvant: Montanide (ISA 51)

Species (Isotype) human (IgG)

**Ab Type** Tat basic region **References** Noonan *et al.* 2003

**Keywords** immunotherapy, vaccine-specific epitope characteristics

· Intramuscular injection of Tat-toxoid induced high titers of anti-Tat reactivity in serum samples of six HIV-1 positive and of four HIV negative study subjects. Anti-Tat Abs successfully blocked extracellular Tat from transactivating HIV Tat-sensitive promotors. The anti-Tat IgG response in sera from two healthy and HIV infected patients inhibited cell entry of synthetic Tat, thus blocking its functional activity. Additionally, the anti-Tat Abs inhibited intercellular Tat transfer in a co-culture cell system. All HIV-1 infected patients had Ab responses to the N-term region of Tat, and 4/4 HIV-1 + and 5/6 HIV-1 negative patients responded to the basic domain. Several additional peptides were recognized either exclusively or more commonly in the HIV+ people. The basic region of Tat mediates binding to VEGFR2 on Kaposi's sarcoma cells and endothelial cells, and HIV patients with Kaposi's sarcoma lack Abs to this domain. Noonan et al. [2003] (vaccine-specific epitope characteristics, immunotherapy)

No. 277

MAb ID 1D2F11

HXB2 Location Tat (49–86)

Author Location Tat

**Epitope** RKKRRQRRRPPQGSQTHQVSLSKQPTSQSRGD-PTGPKE

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Tat

Species (Isotype) mouse (IgG1)

Ab Type C-term

References Valvatne et al. 1996

1D2F11: MAb did not bind shorter peptides – this MAb inhibited exogenously delivered Tat transactivation of an HIV-LTR-CAT construct in HeLa cells by inhibition of cellular uptake of Tat. Valvatne et al. [1996]

**No.** 278

MAb ID 2D9E7

HXB2 Location Tat (49–86)

**Author Location** Tat

Epitope RKKRRQRRRPPQGSQTHQVSLSKQPTSQSRGD-

**PTGPKE** 

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Tat

Species (Isotype) mouse (IgG1)

Ab Type C-term

References Valvatne et al. 1996

2D9E7: MAb did not bind shorter peptides – this MAb inhibited exogenously delivered Tat transactivation of an HIV-LTR-CAT construct in HeLa cells by inhibition of cellular uptake of Tat, but less efficiently than MAbs 1D2F11 or 4B4C4. Valvatne et al. [1996]

**No.** 279

MAb ID 4B4C4 (4B4)

HXB2 Location Tat (49–86)

**Author Location** Tat

**Epitope** RKKRRQRRRPPQGSQTHQVSLSKQPTSQSRGD-PTGPKE

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Tat

Species (Isotype) mouse (IgG1)

Ab Type C-term

References Jensen et al. 1997; Valvatne et al. 1996

• 4B4C4: MAb did not bind shorter peptides – this MAb inhibited exogenously delivered Tat transactivation of an HIV-LTR-CAT construct in HeLa cells by inhibition of cellular uptake of Tat. Valvatne *et al.* [1996]

**No.** 280

MAb ID 5G7D8

HXB2 Location Tat (49–86)

**Author Location** Tat

Epitope RKKRRQRRRPPQGSQTHQVSLSKQPTSQSRGD-

PTGPKE

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Tat

Species (Isotype) mouse (IgG1)

Ab Type C-term

References Valvatne et al. 1996

• 5G7D8: MAb did not bind shorter peptides – this MAb inhibited exogenously delivered Tat transactivation of an HIV-LTR-CAT construct in HeLa cells by inhibition of cellular uptake of Tat, but less efficiently than 1D2F11 or 4B4C4. Valvatne *et al.* [1996]

No. 281

MAb ID polyclonal

HXB2 Location Tat (73-86)

Author Location Tat (73-86 IIIB BH10)

Epitope PTSQPRGDPTGPKE?

Subtype B

Neutralizing

HIV Antibodies Tables Tat Antibodies

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA), macrophage activating lipopeptide-2 (MALP-2)

**Species (Isotype)** mouse (IgA, IgG) **References** Borsutzky *et al.* 2003

**Keywords** adjuvant comparison, genital and mucosal immunity, immunodominance, mucosal immunity, Th1, Th2, vaccine-induced epitopes

• Intranasal immunization of BALB/c mice Tat with MALP-2 induced stronger immune responses than i.p. vaccination of Tat with IFA. Also i.n. MALP-2 vaccination favored Th1 responses, while i.p IFA favored Th2. Ab, T help and CTL responses were observed. MALP-2 enhanced Tat-specific IgA responses in the lung and in the genital tract. Three linear Ab epitopes were recognized. The most frequent response (80% of vaginal lavage, 100% of sera and lung lavage) was to Tat peptide 1-20. The second strongest to Tat 46-60 (20% of vaginal lavage, 50% of lung lavage, 50% of sera). Finally, 50% of the sera of Tat+ IFA i.p. immunized mice recognized Tat 73-86. Borsutzky *et al.* [2003] (adjuvant comparison, genital and mucosal immunity, vaccine-induced epitopes, immunodominance, mucosal immunity, Th1, Th2)

No. 282

MAb ID NT2/4D5.24

HXB2 Location Tat (73–86)

**Author Location** Tat

Epitope PTSQPRGDPTGPKE

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: Tat

Species (Isotype) mouse

Ab Type C-term

References Dingwall et al. 1989

• NT2/4D5.24: Immunoprecipitates and immunoblots HIV-1 tat protein. Dingwall *et al.* [1989]

No. 283

MAb ID polyclonal

HXB2 Location Tat (76-89)

Author Location Tat (76–90 89.6)

Epitope QPRGDPTGPKQKKK

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6, B clade HXBc2 HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA)

Species (Isotype) macaque (IgG)

Ab Type C-term, N-term, Tat basic region

References Silvera et al. 2002

**Keywords** antibody binding site definition and exposure, vaccine antigen design

 Anti-Tat and Tat toxoid responses were raised in rhesus macaques using HxBc2 and 89.6P Tat, and Tat toxoids. High anti-Tat IgG responses were observed in all animals, and helper responses were detected in 8/16, and IFN gamma CTL in 11/16. Ab and proliferative responses were observed, and the trucated 86 amino acid IIIB Tat was more immunogenic than the full 102 amino acid Tat. Tat vaccinated animals when challenged with SHIV89.6 were infected despite a robust immune response. The most consistent reactions to the vaccinations were to peptides in regions: N-term, 1-15 (MEPVDRPLEP-WKHPG), basic domain 46-60 (SYGRKKRQRRAHQ), and 61-91, particularly C-term 76-90 (QPRGDPTGPKQKKK). Silvera *et al.* [2002] (antibody binding site definition and exposure, vaccine antigen design)

No. 284

MAb ID

HXB2 Location Tat

**Author Location** Tat

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: Tat Adjuvant: Incomplete Freund's Adjuvant (IFA), Montanide (ISA 51)

Species (Isotype) human

References Gringeri et al. 1998

**Keywords** immunotherapy

• 14 HIV-1 infected individuals were vaccinated with inactivated Tat (called Tat-toxoid), with the intent of enhancing Tat Ab levels to suppress the negative impact of secreted Tat on immune function. Tat vaccinations were safe and patients developed increased levels of Tat-specific Abs; some patients had increased Tat-specific proliferative responses. CD4 T cells tended to increase a small but significant amount after immunization, and in several patients viral load decreased. Gringeri *et al.* [1998] (immunotherapy)

No. 285

MAb ID L-anti-Tat

HXB2 Location Tat

**Author Location** Tat

**Epitope** 

**Neutralizing** L P (when lipidated)

Immunogen vaccine

Vector/Type: protein HIV component: Tat

Species (Isotype) mouse (IgG1)

Research Contact AGMED, Inc., Bedford, MA USA

References Cruikshank et al. 1997

• L-anti-Tat: Lipidated antibody can be taken up by cells and effectively block IIIB and primary virus HIV-1 replication in actively and latently infected cells. Cruikshank *et al.* [1997]

No. 286

MAb ID polyclonal

HXB2 Location Tat Author Location Tat

Epitope

Subtype A, B, C, CRF01\_AE, D

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Belliard et al. 2003

Tat Antibodies HIV Antibodies Tables

Country France

Keywords rate of progression, subtype comparisons

• Sera from 20 HIV-1 positive individuals were tested for their ability to react with Tat proteins from different clades, and were found to react with subtype A, B, and D, but not with subtype C or CRF01 (AE). Sera from 101 slow progressors and 42 fast progressors were tested for responses to Tat peptides, and compared to responses to gp41 peptide, as anti-Tat antibodies have been shown by others to be elevated in slow progressors. In this study, overall levels of Tat antibodies were not different in the two groups, however relative levels of antibodies to different Tat and gp41 peptides were observed. Belliard *et al.* [2003] (subtype comparisons, rate of progression)

**No.** 287

MAb ID polyclonal

**HXB2 Location** Tat

**Author Location** Tat

**Epitope** 

Neutralizing yes

Immunogen vaccine

Vector/Type: protein HIV component: Tat Adjuvant: Complete Freund's Adjuvant

(CFA), red blood cells

Species (Isotype) mouse (IgG1, IgG2a, IgG3)

References Dominici et al. 2003

**Keywords** adjuvant comparison, immunotherapy, Th1,

• BALB/c mice were immunized intra-peritineally with Tat protein bound to red blood cells via biotin-avidin conjugation. This antigen delivery system was successfully internalized by dendritic cells, and induced more consistent anti-Tat NAbs responses and sightly increased Tat-specific CTL responses relative to Tat protein with CFA. RBC-Tat immunization induced Th1 (IgG2a)and Th2 (IgG1 and IgG3)type immune responses. (adjuvant comparison, immunotherapy, Th1, Th2)

No. 288

MAb ID polyclonal

**HXB2 Location** Tat

**Author Location** Tat

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: chitosan nanoparticles HIV component: Tat Adjuvant: adjuvant oily

structure (IMS)

Species (Isotype) mouse (IgA, IgG)

References Le Buanec et al. 2001

Keywords adjuvant comparison, mucosal immunity

Mice were immunized with Tat toxoid (Tat detoxified by carboxamidation) either intranasally or orally using either adjuvant oily structure (IMS), nanoparticles of chitosan, or microparticles of polylactide-co-glycolide. Each of these strategies triggered IgG and IgA that inhibited Tat activity. Le Buanec et al. [2001] (adjuvant comparison, mucosal immunity)

No. 289

MAb ID polyclonal

**HXB2 Location** Tat

Author Location Tat (IIIB)

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: Tat Adjuvant: Cholera toxin (CT), E. coli mutant heat labile enterotoxin (LT-R72), E. coli heat labile enterotoxin

**Species (Isotype)** mouse (IgG)

References Marinaro et al. 2003

Keywords adjuvant comparison, mucosal immunity

• Intranasal immunization of BALB/c mice with Tat and e.coli heat-labile enterotoxin (LT) and non-toxic LT-R72 LT induced strong antigen-specific IgG Abs which remained stable for one year. Tat-specific IgA reponses were measured in vaginal and intestinal secretions. Immunization of BALB/c mice with native Tat (aa1-86) induced serum IgG directed against an immunodominant epitope (aa1-20) and against a second epitope (aa 46-60). CTL responses were also observed. Anti-Tat serum Abs neutralized Tat activity in a dose-independent manner. C57BL/6 remained unresponsive to Tat immunizations when Tat was co-adminstered with LT or cholera toxin (CT) as adjuvant; BALB/c mice are H-2d, C57BL/6 are H-2b. Congenic BALB.C mice that express H-2b rather than H-2d also could not respond to Tat, suggesting the repsonse to Tat is constrained by the haplotype. Marinaro et al. [2003] (adjuvant comparison, mucosal immunity)

No. 290

MAb ID polyclonal

HXB2 Location Tat

**Author Location** Tat

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein, vaccinia Strain: B clade MN HIV component: gp160, Tat Adjuvant: Incomplete Freund's Adjuvant (IFA), polyphosphazene

Species (Isotype) macaque (IgG)

**References** Pauza et al. 2000

• 16 Macaques mulatta were immunized with Tat toxoid, or with Tat plus gp160, and challenged with the SHIV 89.6PD isolate. Sera from 14/16 animals neutralized Tat *in vitro*. 8 macaques developed both cellular and humoral responses to Tat, and 7/8 of these had low viral set points after rectal challenge with SHIV89.6PD. CD4+ T cells in Tat vaccinated infected animals had lower IFN-alpha and chemokine receptor expression, features of infection associated with extracellular Tat. Pauza *et al.* [2000]

**No.** 291

MAb ID polyclonal

**HXB2 Location** Tat

**Author Location** Tat

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

**HIV Antibodies Tables** Tat Antibodies

> *Vector/Type:* protein *Strain:* B clade BH10 HIV component: gp120, Nef, Tat Adjuvant: AS02A (oil-in-water emulsion, 3Dmonophosphoryl lipid A, QS21), AS06 (CpG, Author Location Tat (IIIB, 89.6, CMU08) aluminum hydroxide)

Species (Isotype) macaque (IgG) References Voss et al. 2003

> Keywords adjuvant comparison, variant crossrecognition or cross-neutralization

· Monkeys were immunized with different combinations of gp120/NefTat/SIV-Tat three times over a three-month time period and intravenously challenged with SHIV 89.6P four weeks after the last immunzation. Vaccination induced anti-Tat, -Nef and -gp120 Ab responses that decreased until challenge. Neutralizing Ab responses against HIV-1 MN and HIV-1 W61D were observed. Vaccination with the combination of all three proteins reduced viral load and kept animals from a CD4+ T cell decline, preventing AIDS for more than 2.5 years. The adjuvant AS02A elicited NAbs and protected from disease, while AS06 did not. gp120 alone or TatNef/SIV-Tat without gp120 could not protect from disease. Voss et al. [2003] (adjuvant comparison, variant cross-recognition or crossneutralization)

**No.** 292

MAb ID polyclonal

**HXB2 Location** Tat

Author Location Tat

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Zagury et al. 1998

Country France

Keywords immunotherapy, rate of progression

• Comparing 67 fast progressors with 182 non-progressors in the GRIV cohort, only anti-Tat Ab levels, not Abs to Env, Gag, or Nef, were correlated as a serological indicator of rate of progression. This suggests that raising Tat Abs may be beneficial as immunotherapy or in a vaccine. Zagury et al. [1998] (immunotherapy, rate of progression)

**No.** 293

MAb ID 2D9D5

**HXB2 Location** Tat

Author Location Tat

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Tat

Species (Isotype) mouse (IgG)

Ab Type C-term

References Mhashilkar et al. 1995

• 2D9D5: Single chain antibodies, intrabodies, were engineered that can be stably expressed in the cytoplasm of mammalian cells - co-expression of C-term intrabody did not inhibit transactivation of an HIV LTR-CAT construct, in contrast to MAb 1D9D5. Mhashilkar et al. [1995]

No. 294

MAb ID polyclonal

**HXB2 Location** Tat

**Epitope** 

Subtype B, CRF01\_AE

**Neutralizing** 

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade HIV

component: Tat

Species (Isotype) human (IgG)

Ab Type C-term, N-term, Tat basic region

References Richardson et al. 2003

**Country** France

Keywords antibody binding site definition and exposure, rate of progression, subtype comparisons, vaccine-specific epitope characteristics

• Serum samples were obtained from the French GRIV (genetic resistance to HIV) cohort and tested for anti-Tat and anti-Vpr responses as both may contribute as extracellular proteins to pathogenesis. Serum anti-Tat IgG responses were significantly higher and maintained for up to 20 months in stable non-progressors (CD4+ T cell counts greater than 500 cell/ul after being positive for 8 years with no ART) compared to unstable non-progressors and fast progressors (CD4+ T less than 300 cells/ul within 2 years of seroconversion, some HAART). Anti-Tat IgG from GRIV stable non-progressors recognized linear epitopes located within the N-terminal, basic and the Cterminal domains of Tat. Humoral responses of fast-progressors and of one unstable non-progressor were restricted to the basic region of Tat. Tat toxoid vaccinees from Milan tended to recognize N-terminal and C-terminal domains. Sera from some GRIV and Tat toxoid vaccinees cross-reacted in an ELISA assay with a truncated 89.6 S/HIV 89.6P Tat, 89.6P Tat, HIv-1 subtype E (CMU08) and with SIVmac251 Tat (one sample). Richardson et al. [2003] (antibody binding site definition and exposure, vaccine-specific epitope characteristics, subtype comparisons, rate of progression)

No. 295

MAb ID polyclonal

**HXB2 Location** Tat

**Author Location** Tat

**Epitope** 

Subtype B, CRF01\_AE

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6,

B clade IIIB HIV component: Tat

**Species (Isotype)** macaque (IgG)

Ab Type C-term, N-term, Tat basic region

References Richardson et al. 2002

Keywords antibody binding site definition and exposure, vaccine antigen design, variant crossrecognition or cross-neutralization

 Anti-Tat responses were raised in rhesus macaques using IIIB Tat, SHIV89.6P Tat, carboxymethylated Tat and 89.6P Tat toxoids. Tat IgG responses to the vaccine were cross-reactive with subtype E and MAC 251. Ab and proliferative responses were observed, and the trucated 86 amino acid IIIB Tat was more Tat Antibodies HIV Antibodies Tables

immunogenic than the full 102 amino acid Tat. Tat vaccinated animals when challenged with SHIV89.6 were infected despite a robust immune response and were not distinguishable from controls. Richardson *et al.* [2002] (antibody binding site definition and exposure, vaccine antigen design, variant cross-recognition or cross-neutralization)

No. 296 MAb ID polyclonal HXB2 Location Tat

Author Location Tat (IIIB, 89.6, CMU08)

**Epitope** 

Subtype B, CRF01\_AE

Neutralizing

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade HIV

component: Tat

Species (Isotype) human (IgG)

Ab Type C-term, N-term, Tat basic region

References Richardson et al. 2003

Country France

**Keywords** antibody binding site definition and exposure, rate of progression, subtype comparisons, vaccine-specific epitope characteristics

• Serum samples were obtained from the French GRIV (genetic resistance to HIV) cohort and tested for anti-Tat and anti-Vpr responses as both may contribute as extracellular proteins to pathogenesis. Serum anti-Tat IgG responses were significantly higher and maintained for up to 20 months in stable non-progressors (CD4+ T cell counts greater than 500 cell/ul after being positive for 8 years with no ART) compared to unstable non-progressors and fast progressors (CD4+ T less than 300 cells/ul within 2 years of seroconversion, some HAART). Anti-Tat IgG from GRIV stable non-progressors recognized linear epitopes located within the N-terminal, basic and the Cterminal domains of Tat. Humoral responses of fast-progressors and of one unstable non-progressor were restricted to the basic region of Tat. Tat toxoid vaccinees from Milan tended to recognize N-terminal and C-terminal domains. Sera from some GRIV and Tat toxoid vaccinees cross-reacted in an ELISA assay with a truncated 89.6 S/HIV 89.6P Tat, 89.6P Tat, HIv-1 subtype E (CMU08) and with SIVmac251 Tat (one sample). Richardson et al. [2003] (antibody binding site definition and exposure, vaccine-specific epitope characteristics, subtype comparisons, rate of progression)

No. 297
MAb ID G1
HXB2 Location Tat
Author Location Tat (1–15)
Epitope
Subtype B
Neutralizing yes

Strain: B clade HIV component: Tat

**Species (Isotype)** human (IgG1 $\kappa$ )

Immunogen vaccine

Ab Type N-term

References Moreau et al. 2004

**Keywords** antibody binding site definition and exposure, antibody sequence, variable domain, subtype comparisons

• G1: G1 is a single-chain fragment-variable scFv antibody derived from a Tat-toxoid vaccinated uninfected volunteer. G1 binds strongly to soluble rTAT protein and to denatured rTAT, suggesting that the epitope is linear. G1 recognized HIV-1 clade B Tat proteins Bru and HXB2, but did not bind to clade E (CM240), clade C (92Br), clade D (Eli) and clade A (Ug11RP). G1 inhibited Tat-transactivation of viral replication. The VH3 heavy chains of the two phage scFvG1 VH3 heavy chain sequences of scFvG1 and scFvG2 vary (G1 CDR3, RGSTGKALDYCSPRTL; G2 CDR3, ERSQQHCN-PLLHSNGKNYAE) although both share identical Vk light chain sequences. Moreau et al. [2004] (antibody binding site definition and exposure, subtype comparisons, antibody sequence, variable domain)

No. 298

MAb ID G2

**HXB2 Location** Tat

**Author Location** Tat (1–15)

Epitope
Subtype B
Neutralizing yes
Immunogen vaccine

Strain: B clade HIV component: Tat

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type N-term

References Moreau et al. 2004

**Keywords** antibody binding site definition and exposure, antibody sequence, variable domain, subtype comparisons

• G2: G2 is a single-chain fragment-variable scFv antibody derived from a Tat-toxoid vaccinated uninfected volunteer. G2 binds strongly to soluble rTAT protein and to denatured rTAT, suggesting that the epitope is linear. G2 recognized HIV-1 clade B Tat proteins Bru and HXB2, but did not bind to clade E (CM240), clade C (92Br), clade D (Eli) and clade A (Ug11RP). G2 inhibited Tat-transactivation of viral replication. The VH3 heavy chains of the two phage scFvG1 VH3 heavy chain sequences of scFvG1 and scFvG2 vary (G1 CDR3, RGSTGKALDYCSPRTL; G2 CDR3, ERSQQHCN-PLLHSNGKNYAE) although both share identical Vk light chain sequences and Tat binding sites. Moreau et al. [2004] (antibody binding site definition and exposure, subtype comparisons, antibody sequence, variable domain)

No. 299

MAb ID J1

HXB2 Location Tat

Author Location Tat (1-15)

**Epitope** 

Subtype B

**Neutralizing** yes

Immunogen vaccine

Strain: B clade HIV component: Tat

**Species (Isotype)** human (IgG1 $\lambda$ )

**Ab Type** N-term

References Moreau et al. 2004

HIV Antibodies Tables Tat Antibodies

**Keywords** antibody binding site definition and exposure, antibody sequence, variable domain, subtype comparisons

• J1: J1 is a single-chain fragment-variable scFv antibody derived from a Tat-toxoid vaccinated uninfected volunteer. J1 binds strongly to soluble rTAT protein and to denatured rTAT, suggesting that the epitope is linear. J1 recognized HIV-1 clade B Tat proteins Bru and HXB2, but did not bind to clade E (CM240), clade C (92Br), clade D (Eli) and clade A (Ug11RP). J1 inhibited Tat-transactivation of viral replication. Of three scFv antibodies, all bound the N-terminal amino acids 1-15, but G1 and G2 had kappa light chains and J1 had lambda, and the CDR3 of each was distinct, with J1's CDR3 sequence being: RDRYC-SSPGCYKGADGGRLKDY. Moreau et al. [2004] (antibody binding site definition and exposure, subtype comparisons, antibody sequence, variable domain)

No. 300 MAb ID TC15 HXB2 Location Tat

Author Location Tat (Lai/Bru)

Epitope
Subtype B
Neutralizing
Immunogen vaccine

Vector/Type: protein Strain: B clade BRU
HIV component: Tat Adjuvant: Complete

Freund's Adjuvant (CFA)

 $\boldsymbol{Species}\;(\boldsymbol{Isotype})\;\;\text{mouse}\;(\boldsymbol{IgG})$ 

Ab Type N-term

Research Contact Dr. J.-L. Guesdon, Institut Pasteur, Paris

**References** Belliard *et al.* 2003 **Keywords** subtype comparisons

• TC15: This is one of 5 anti-Tat murine monoclonal antibodies generated in this study. It is conformational reacting only with intact protein. It reacts with B and D clade Tat proteins, and does not recognize Tat from clade A, C, or CRF01 (AE). Belliard *et al.* [2003] (**subtype comparisons**)

**No.** 301

MAb ID polyclonal

**HXB2 Location** Tat

Author Location Tat (Lai/Bru)

Epitope Subtype B Neutralizing

Immunogen SHIV infection, vaccine

Vector/Type: peptide Strain: B clade BRU HIV component: Tat Adjuvant: aluminum phosphate, CpG immunostimulatory sequence (ISS), Montanide (ISA 720)

Species (Isotype) macaque (IgG)

Ab Type N-term

References Belliard et al. 2003

**Keywords** rate of progression

 Macaques were immunized with different combinations of Tat peptides. Serum from these animals was able to inhibit Tat-induced apoptosis, and Tat antibodies are associated with long term survival. Anti-Tat antibodies generated in infected macaques tended to be restricted to the peptide 44-61, while sera from infected humans could react with several different peptides. Belliard *et al.* [2003] (**rate of progression**)

**No.** 302

MAb ID polyclonal

**HXB2 Location** Tat

Author Location Tat (Lai/Bru)

Epitope Subtype B Neutralizing

Immunogen SHIV infection, vaccine

Vector/Type: peptide Strain: B clade BRU HIV component: Tat Adjuvant: BSA, Com-

plete Freund's Adjuvant (CFA)

Species (Isotype) rabbit (IgG)

Ab Type N-term

**References** Belliard *et al.* 2003 **Keywords** rate of progression

• 12 rabbits were immunized with different combinations of Tat peptides. Abs raised against peptide aa 8-53 did not react with the peptide 19-53, suggesting that the N-terminal region is important. Serum from these animals was able to inhibit Tat-induced apoptosis, and Tat antibodies in humans are associated with long term survival. Belliard *et al.* [2003] (**rate of progression**)

No. 303

MAb ID B1E3

**HXB2 Location** Tat

**Author Location** Tat (44–61)

Epitope
Subtype B
Neutralizing yes
Immunogen vaccine

Strain: B clade HIV component: Tat

**Species (Isotype)** human (IgG1 $\kappa$ ) **Ab Type** Tat basic region

References Moreau *et al.* 2004

**Keywords** antibody binding site definition and exposure, subtype comparisons

• B1E3: B1E3 is a MAb derived from a Tat-toxoid vaccinated uninfected volunteer. B1E3 recognized two Tat peptides, aa19-53 and aa44-61 of an unspecified HIV-1 clade B Tat protein. B1E3 demonstrates a weak binding affinity to rTAT protein in solution, suggesting that epitope recognition may be conformation dependent B1E3 did not recognize synthetic HIV-1 clade B Tat proteins Bru and HXB2, clade E (CM240), clade C (92Br), clade D (Eli) and clade A (Ug11RP). It only bound to native TAT protein, and could inhibit Tat-transactivation. Moreau et al. [2004] (antibody binding site definition and exposure, subtype comparisons)

**No.** 304

MAb ID J3B2

**HXB2 Location** Tat

**Author Location** Tat (44–61)

Epitope

Subtype B

Neutralizing yes

Immunogen vaccine

**Rev Antibodies HIV Antibodies Tables** 

Strain: B clade HIV component: Tat

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type Tat basic region References Moreau et al. 2004

subtype comparisons

• J3B2: J3B2 is a MAb derived from a Tat-toxoid vaccinated uninfected volunteer. B1E3 recognized two Tat peptides, aa33-37 and aa37-51 of an unspecified HIV-1 clade B Tat protein. J3B2 demonstrates a weak binding affinity to rTAT protein in solution, suggesting that epitope recognition may be conformation dependent, B1E3 did not recognize synthetic HIV-1 clade B Tat proteins Bru and HXB2, clade E (CM240), clade C (92Br), clade D (Eli) and clade A (Ug11RP). It only bound to native TAT protein, and could inhibit Tat-transactivation. Moreau et al. [2004] (antibody binding site definition and exposure, subtype comparisons)

## **IV-C-14** Rev Antibodies

**No.** 305

MAb ID 4G9

HXB2 Location Rev (5-15)

**Author Location** Rev (5–15)

Epitope SGDSDEELIRT?

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Rev

Species (Isotype) mouse

References Jensen et al. 1997

• 4G9: Mapped binding location by protein footprinting. Jensen et al. [1997]

**No.** 306

MAb ID Ab2

HXB2 Location Rev (32-50)

Author Location Rev (32-49 BRU)

Epitope EGTRQARRNRRRWRERQR

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Rev

Species (Isotype) (IgG1)

Research Contact Tony Lowe and Jonathan Karn, MRC Center, Cambridge

**References** Henderson & Percipalle 1997

• Ab2: The Ab2 binding site overlaps the nuclear localization signal - Ab2 binding to Rev was blocked by bound HIV RNA - the cellular protein importin-beta can bind in this Arg rich region – atypically, the Rev binds specifically to importin-beta, but not to the importin-beta-importin-alpha dimer. Henderson & Percipalle [1997]

No. 307

**MAb ID** 10.1

HXB2 Location Rev (33-48)

Author Location Rev (33-48)

Epitope GTRQARRNRRRRWRER?

**Neutralizing** 

**Immunogen** 

Species (Isotype)

References Maksiutov et al. 2002; Ranki et al. 1995; Ranki et al. 1994; Ovod et al. 1992

- **Keywords** antibody binding site definition and exposure, 10.1: This epitope is similar to a fragment of the human protein Complement 4 (containing C4A anaphlylatoxin), GRRNRRRR. Maksiutov et al. [2002]
  - 10.1: Binds to the RRE binding site polyclonal anti-Rev Ab detected Rev in astrocytes in 4/5 brain autopsy samples, but only one of these was positive using 10.1, suggesting most Rev was bound to RRE. Ranki et al. [1995]

No. 308

MAb ID 3H6

HXB2 Location Rev (38-43)

Author Location Rev (38-44)

**Epitope** RRNRRR

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Rev

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Maksiutov et al. 2002; Orsini et al. 1995

- 3H6: There is another MAb with this ID that recognizes gp41.
- 3H6: This epitope is similar to a fragment of the human protein Complement 4 (containing C4A anaphlylatoxin), GRRNRRRR. Maksiutov et al. [2002]
- 3H6: Directed against nucleolar localization/RRE binding domain - antigenic domain tentative, MAb failed to bind a RRN-RRR Rev deletion mutant. Orsini et al. [1995]

No. 309

MAb ID 8E7

HXB2 Location Rev (70-84)

Author Location Rev (70-84)

Epitope PVPLQLPPLERLTLD

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Rev

Species (Isotype) mouse (IgG2a $\kappa$ )

References Maksiutov et al. 2002; Boe et al. 1998; Jensen et al. 1997; Szilvay et al. 1995; Kalland et al. 1994b; Kalland et al. 1994a

- 8E7: This epitope is similar fragments of the human protein Epidermal growth factor receptor substrate 15, EPVPMSLPPA, and Insulin-like growth factor binding protein complex acid labile chain precursor, QPPGLERLWLEGNPWDCG. Maksiutov et al. [2002]
- 8E7: HIV-1 RNA and Rev localize to the same region in the nucleoplasm, but the splicing factor SC-35 localizes in different speckles with the nucleoplasm than Rev – intron containing beta-globin was distributed similarly to HIV-1, suggesting Rev and HIV-1 RNAs interact at putative sites of mRNA transcriptions and splicing. Boe et al. [1998]
- 8E7: Peptide interaction mapped to aa 70-84, 75-88 protein footprint to 65-88. Jensen et al. [1997]
- 8E7: 8E7 worked in indirect immunofluorescence and also detected Rev in WB assays - used to detect localization of Rev in several compartments including the nucleoli, nucleoplasm, perinuclear zone, and cytoplasm - Rev co-localized with host cell factors known to assemble on nascent transcripts - Rev

**HIV Antibodies Tables Rev Antibodies** 

shuttles continuously between cytoplasmic and nucleoplasmic compartments. Kalland et al. [1994a,b]; Szilvay et al. [1995]

**No.** 310

MAb ID 9G2 (9G2G4D6E8)

HXB2 Location Rev (70-84)

Author Location Rev (70-84)

Epitope PVPLQLPPLERLTLD

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Rev

**Species (Isotype)** mouse (IgG2a $\kappa$ ) Research Contact Anne Marie Szilvay

> References Maksiutov et al. 2002; Jensen et al. 1997; Research Contact Anne Marie Szilvay Kalland et al. 1994a

- 9G2: Called 9G2G4D6E8: UK Medical Research Council AIDS reagent: ARP3058.
- 9G2: This epitope is similar fragments of the human protein Epidermal growth factor receptor substrate 15, EPVPMSLPPA, and Insulin-like growth factor binding protein complex acid labile chain precursor, QPPGLERLWLEGNPWDCG. Maksiutov et al. [2002]
- 9G2: Peptide interaction mapped to aa 70-84, 75-88 protein footprint to 65-88. Jensen et al. [1997]
- 9G2: Worked in indirect immunofluorescence and also detected Rev in WB assays - used to detect localization of Rev throughout the cell. Kalland et al. [1994a]

No. 311

MAb ID Ab4

HXB2 Location Rev (72–91)

**Author Location** Rev (72–91 BRU)

Epitope PLQLPPLERLTLDCNEDCGT

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Rev

Species (Isotype) (IgG1)

Research Contact Tony Lowe and Jonathan Karn, MRC Center,

Cambridge

References Maksiutov et al. 2002; Henderson & Percipalle 1997

- Ab4: This epitope is similar fragments of the human protein Epidermal growth factor receptor substrate 15, EPVPMSLPPA, and Insulin-like growth factor binding protein complex acid labile chain precursor, QPPGLERLWLEGNPWDCG. Maksiutov et al. [2002]
- Ab4: The binding site overlaps the nuclear export signal binding was not blocked by bound HIV RNA and may be accessible for protein interaction. Henderson & Percipalle [1997]

**No.** 312

MAb ID 3G4

HXB2 Location Rev (90-116)

Author Location Rev (90-116)

Epitope GTSGTQGVGSPQILVESPTVLESGTKE?

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Rev

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

References Orsini et al. 1995

• 3G4: Binds to a region that can be dispensed with and still retain Rev function. Orsini et al. [1995]

**No.** 313

**MAb ID** 1G10 (IG10F4)

HXB2 Location Rev (96–105)

Author Location Rev (95-105)

Epitope GVGSPQILVE

**Neutralizing** 

Immunogen vaccine

*Vector/Type:* protein *HIV component:* Rev

**Species (Isotype)** mouse (IgG2b $\kappa$ )

References Jensen et al. 1997: Kalland et al. 1994a

- 1G10: Called IG10F4: UK Medical Research Council AIDS reagent: ARP3060.
- 1G10: Peptide interaction mapped to aa 91-105, 96-110 protein footprint to aa 10-20, and 95-105. Jensen et al. [1997]
- 1G10: Bound Rev in indirect immunofluorescence and also detected Rev in WB - used to detect localization of Rev throughout the cell. Kalland et al. [1994a]

No. 314

MAb ID 1G7

HXB2 Location Rev (96-105)

Author Location Rev (95-105)

Epitope GVGSPQILVE

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Rev

**Species (Isotype)** mouse (IgG2b $\kappa$ )

References Jensen et al. 1997; Kalland et al. 1994a

- 1G7: Peptide interaction mapped to aa 91-105, 96-110 protein footprint to aa 95-105. Jensen et al. [1997]
- · 1G7: Worked in indirect immunofluorescence and also detected Rev in WB – used to detect localization of Rev throughout the cell. Kalland et al. [1994a]

No. 315

MAb ID Ab3

HXB2 Location Rev (102–116)

Author Location Rev (102-116 BRU)

Epitope ILVESPTVLESDKTE

**Neutralizing** 

Immunogen vaccine

*Vector/Type:* protein *HIV component:* Rev

Species (Isotype) (IgG1)

Research Contact Tony Lowe and Jonathan Karn, MRC, Cambridge

References Henderson & Percipalle 1997

• Ab3: This binding site is at the carboxy end of Rev - Ab3 binding was not blocked by bound HIV RNA. Henderson & Percipalle [1997]

No. 316

MAb ID 2G2

**HXB2 Location** Rev

**Author Location Rev** 

**Epitope** 

Vpu Antibodies **HIV Antibodies Tables** 

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Rev

**Species (Isotype)** mouse (IgG1 $\kappa$ ) References Orsini et al. 1995

• 2G2: Does not bind to any of a set of glutathione S-transferase (GST) Rev fusion proteins, or to Rev in a RIPA buffer, suggesting a conformational epitope. Orsini et al. [1995]

## IV-C-15 Vpu Antibodies

**No.** 317

MAb ID DE7

HXB2 Location Vpu (42-63)

Author Location Vpu (41-62)

Epitope LIDRLIERAEDSGNESEGEISA

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: Vpu

Adjuvant: BSA

Species (Isotype) mouse (IgG1)

References Gharbi-Benarous et al. 2004

Keywords antibody binding site definition and exposure,

antibody generation

• DE7: This MAb was generated against the phosphorylated Vpu41-62 peptide. Phosphorylation of Vpu, at the Serines of the DSGXXS motif, is required for interaction of Vpu with the ubiquitin ligase that triggers CD4 degradation and infectious virion release. DE7 bound peptide conformation was analyzed using STD NMR epitope mapping, TRNOESY conformational analysis and molecular dynamics simulation, and found to adopt a compact structure with several bends, including a tight bend at DpSGNEpS. Gharbi-Benarous et al. [2004] (antibody binding site definition and exposure, antibody generation)

# IV-C-16 gp160 Antibodies

**No.** 318

MAb ID M85

**HXB2 Location** gp160 (30–51)

**Author Location** gp120 (30–51 LAI)

Epitope ATEKLWVTVYYGVPVWKEATTT

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) mouse (IgG1)

Ab Type gp120 C1

Research Contact Fulvia di Marzo Veronese

References Koefoed et al. 2005; Wyatt et al. 1997; Ditzel et al. 1997: Moore & Sodroski 1996: Moore

et al. 1994d; Moore et al. 1994c; di Marzo

Veronese et al. 1992

**Keywords** antibody binding site definition and exposure

• M85: IgG antibody phage display libraries were created from HIV-1 + individuals after pre-selection of PBMC with gp120, as an alternative to using bone marrow for generating libraries. M85 was among a set of Abs used for competition studies to define the binding sites of the newly isolated MAbs, and has a linear C1 epitope. Koefoed et al. [2005] (antibody binding site definition and exposure)

- M85: Binds efficiently to sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding. Wyatt et al. [1997]
- M85: Binding inhibited by MAb 4D4#85, enhanced by conformationally sensitive anti-V3 MAb 5G11, and some anti-18 MAbs. Moore & Sodroski [1996]
- M85: C1 domain mutation 40 Y/D impairs binding the relative affinity for denatured/native gp120 is < .01, suggesting conformational component. Moore et al. [1994c]
- M85: Immunoblot and RIP reactive for strains IIIB, 451, MN, RF, and RUTZ – binds deglycosylated gp120. di Marzo Veronese *et al.* [1992]

**No.** 319

**MAb ID** 7E2/4

**HXB2 Location** gp160 (31–50)

Author Location gp120 (31–50 LAI)

**Epitope** TEKLWVTVYYGVPVWKEATT

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

Research Contact S. Ranjbar, NIBSC, UK

References Maksiutov et al. 2002; Moore et al. 1994c

- 7E2/4: UK Medical Research Council AIDS reagent: ARP3050.
- 7E2/4: This epitope has a high degree of similarity with the platelet membrane glycoprotein IIIA precursor (GLIIIA) (integrin beta- 3) (CD61): PLYKEATSTF. Maksiutov et al. [2002]
- 7E2/4: C1 domain the relative affinity for denatured/native gp120 is .07, suggesting conformational component. Moore et al. [1994c]

No. 320

MAb ID 4D4#85

**HXB2 Location** gp160 (41–50)

Author Location gp120 (LAI)

**Epitope** GVPVWKEATT

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

Research Contact S. Nigida and L. Arthur, NCI, Frederick, MD

USA

References Maksiutov et al. 2002; Binley et al. 1998;

Wyatt et al. 1997; Moore & Sodroski 1996; Moore et al. 1994d; Moore et al. 1994c

- 4D4#85: This epitope has a high degree of similarity with the platelet membrane glycoprotein IIIA precursor (GLIIIA) (integrin beta- 3) (CD61): PLYKEATSTF. Maksiutov *et al.* [2002]
- 4D4#85: A panel of MAbs were shown to bind with similar
  or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta
  V1, V2, and V3), thus such a core protein produces a structure
  closely approximating full length folded monomer. Binley et al.
  [1998]
- 4D4#85: Binds efficiently to sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding does not bind to HXBc2 gp120 if the 19 C-term amino acids, in conjunction with C1 positions 31-50, are deleted. Wyatt *et al.* [1997]
- 4D4#85: Inhibits binding of C1 MAb M85, C1-C5 discontinuous epitope MAbs 181 and 212A, and CD4 binding induced MAbs 48d and 17b. Moore & Sodroski [1996]
- 4D4#85: C1 domain the relative affinity, denatured/native gp120 is 0.1 mutation 45 W/S impairs binding. Moore *et al.* [1994c]

**No.** 321

MAb ID M92

**HXB2 Location** gp160 (41–50)

Author Location gp120 (31-50 LAI)

Epitope GVPVWKEATT

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) rat (IgG1)

Ab Type gp120 C1

Research Contact Fulvia di Marzo Veronese

**References** Maksiutov *et al.* 2002; Moore *et al.* 1994d; Moore *et al.* 1994c; di Marzo Veronese *et al.* 1992

- M92: This epitope has a high degree of similarity with the platelet membrane glycoprotein IIIA precursor (GLIIIA) (integrin beta- 3) (CD61): PLYKEATSTF. Maksiutov et al. [2002]
- M92: The relative affinity for denatured/native gp120 is 1. Moore *et al.* [1994c]
- M92: Immunoblot reactive, RIP negative, but precipitates deglycosylated gp120 reacts with strains IIIB, 451, MN, RF, and RUTZ. di Marzo Veronese et al. [1992]

No. 322

MAb ID M86

**HXB2 Location** gp160 (42–61)

**Author Location** gp120 (42–61 LAI)

Epitope VPVWKEATTTLFCASDAKAY

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) mouse (IgG1)

Ab Type gp120 C1

Research Contact Fulvia di Marzo Veronese

**References** Maksiutov *et al.* 2002; Moore *et al.* 1994c; di Marzo Veronese *et al.* 1992

 M86: This epitope has a high degree of similarity with the platelet membrane glycoprotein IIIA precursor (GLIIIA) (integrin beta- 3) (CD61): PLYKEATSTF. Maksiutov et al. [2002]

- M86: C1 domain the relative affinity for denatured/native gp120 is 1. Moore *et al.* [1994c]
- M86: Immunoblot and RIP reactive for strains IIIB, 451, MN, RF, and RUTZ – binds deglycosylated gp120. di Marzo Veronese et al. [1992]

No. 323

MAb ID polyclonal

HXB2 Location gp160 (52-71)

Author Location Env (42-61 LAI)

Epitope LFCASDAKAYDTEVHNVWAT

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia HIV component: Env

Species (Isotype) mouse

Ab Type gp120 C1

References Collado et al. 2000

• Vaccinia p14 can elicit NAbs and p39 tends to be immunodominant, so these two proteins were fused to regions of HIV-1 Env – reduced glycosylation was noted when p14 or p39 was placed in the N-term region of the fusion protein – chimeric proteins shifted the Env Ab response from V3 to either a C1 or C4 domain, depending on the construct – all chimeric Env proteins: 14kEnv, 39kEnv, and Env39k elicited a strong Ab response to the C1 region of gp120 (LFCASDAKAYDTEVHNVWAT), and Env39k mounted a strong response to the C4 region (KAMYAPPISGQIRCSSNITG) Collado et al. [2000]

No. 324

**MAb ID** 133/237

HXB2 Location gp160 (61-70)

Author Location gp120 (51-70 LAI)

Epitope YDTEVHNVWA

Subtype B

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

**Species (Isotype)** mouse (IgG1)

Ab Type gp120 C1

**References** Pantophlet *et al.* 2004; Moore *et al.* 1994d; Moore *et al.* 1994c; Niedrig *et al.* 1992b

**Keywords** vaccine antigen design

• 133/237: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including 133/237. Pantophlet *et al.* [2004] (vaccine antigen design)

- mutation of position 69 W/L impairs binding. Moore et al. [1994c]

weak neutralization of lab strains. Niedrig et al. [1992b]

No. 325

MAb ID 133/290

HXB2 Location gp160 (61-70)

Author Location gp120 (61-70 LAI)

Epitope YDTEVHNVWA

Subtype B

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 C1

Research Contact M. Niedrig

References Pantophlet et al. 2003b; Yang et al. 2000; Binley et al. 1998; Wyatt et al. 1997; Binley et al. 1997a; Moore & Sodroski 1996; Wyatt et al. 1995; Moore et al. 1994d; Moore et al. 1994c; Thali et al. 1993; Niedrig et al. 1992b

Keywords vaccine antigen design

- 133/290: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)
- 133/290: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes – CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 – non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 - MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang et al. [2000]
- 133/290: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer. Binley et al.
- 133/290: Binds efficiently to sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding. Wyatt et al. [1997]
- 133/290: Reciprocal binding inhibition with the antibody 522-149, that binds to a discontinuous epitope – binding is enhanced by some C5 and C1 binding site antibodies. Moore & Sodroski [1996]

• 133/237: The relative affinity, denatured/native gp120 is 1.4 • 133/290: Used for antigen capture assay, either to bind gp120 to the ELISA plate, or to quantify bound gp120. Wyatt et al.

- 133/237: Region of overlap for reactive peptides is WATHA • 133/290: The relative affinity for denatured/native gp120 is 2.2 - mutation in position 69 W/L impairs binding. Moore et al. [1994c]
  - 133/290: Region of overlap for reactive peptides is WATHA weak neutralization of lab strains. Niedrig et al. [1992b]

No. 326

MAb ID 133/11

HXB2 Location gp160 (64–78)

Author Location gp120 (64-78)

Epitope EVHNVWATHACVPTD

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 C1

References Niedrig et al. 1992b

• 133/11: Region of overlap for reactive peptides is WATHA weak neutralization of lab strains. Niedrig et al. [1992b]

No. 327

MAb ID D/3G5

HXB2 Location gp160 (73-82)

Author Location gp120 (73–82 LAI)

Epitope ACVPTDPNPQ

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 C1

References Bristow et al. 1994

D/3G5: C1 MAb generated in a study of the humoral immune response to Baculovirus-expressed mis-folded rgp120 and rgp160. Bristow et al. [1994]

No. 328

MAb ID D/6A11

HXB2 Location gp160 (73–82)

Author Location gp120 (73-82 LAI)

Epitope ACVPTDPNPQ

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 C1

References Bristow et al. 1994

• D/6A11: C1 MAb generated in a study of the humoral immune response to Baculovirus-expressed mis-folded rgp120 and rgp160. Bristow et al. [1994]

No. 329

MAb ID D/5E12

**HXB2 Location** gp160 (73–92) Epitope PQEVVLVNVTENFDMWKNDM Author Location gp120 (73-92 LAI) Subtype B Epitope ACVPTDPNPQEVVLVNVTEN Neutralizing L Subtype B Immunogen vaccine Neutralizing no Vector/Type: protein Strain: B clade IIIB HIV component: gp120 Immunogen vaccine Vector/Type: protein Strain: B clade LAI Species (Isotype) rat HIV component: gp120 Ab Type gp120 C1 Species (Isotype) mouse References Moore et al. 1994c; Nakamura et al. 1992; Ab Type gp120 C1 Berman et al. 1991: Dowbenko et al. 1988 References Bristow et al. 1994 • 1D10: The relative affinity for denatured/native gp120 is 13 – • D/5E12: C1 MAb generated in a study of the humoral immutation 88 N/P impairs binding. Moore et al. [1994c] mune response to Baculovirus-expressed mis-folded rgp120 1D10: Cross-blocks 5B3 in IIIB-rsgp160 ELISA – type specific and rgp160. Bristow et al. [1994] in rgp120 ELISA binding. Nakamura et al. [1992] **No.** 330 No. 333 MAb ID L5.1 MAb ID B242 HXB2 Location gp160 (79–93) HXB2 Location gp160 (83-92) Author Location gp120 (89-103 IIIB) Author Location gp120 (83–92 LAI) **Epitope** PNPQEVVLVNVTENF **Epitope** EVVLVNVTEN Neutralizing Subtype B Immunogen vaccine Neutralizing no Vector/Type: vaccinia Strain: B clade IIIB Immunogen vaccine HIV component: gp160 Vector/Type: protein Strain: B clade NL43 Species (Isotype) mouse (IgG) HIV component: gp160 Ab Type gp120 C1 **Species (Isotype)** mouse (IgG1) References Akerblom et al. 1990 Ab Type gp120 C1 References Bristow et al. 1994 No. 331 • B242: C1 MAb generated in a study of the humoral immune re-MAb ID 4A7C6 sponse to Baculovirus-expressed mis-folded rgp160 IIIB:NL43, **HXB2 Location** gp160 (81–90) MicroGenSys. Bristow et al. [1994] Author Location gp120 (81–90 LAI) Epitope PQEVVLVNVT No. 334 Subtype B **MAb ID** 133/192 **Neutralizing** HXB2 Location gp160 (91-100) Author Location gp120 (91–100 LAI) Immunogen vaccine *Vector/Type:* protein *HIV component:* Env **Epitope** ENFDMWKNDM Species (Isotype) mouse (IgG) Subtype B Ab Type gp120 C1 Neutralizing L Research Contact R. Tedder Immunogen vaccine References Moore & Sodroski 1996; Moore et al. 1994d; Vector/Type: protein Strain: B clade IIIB Moore et al. 1994c; Moore & Ho 1993; Thali HIV component: gp120 et al. 1993; Thiriart et al. 1989 **Species (Isotype)** mouse (IgG1)

• 4A7C6: UK Medical Research Council AIDS reagent: ARP

- 4A7C6: Reciprocal binding inhibition with the antibody 133/192 enhanced by anti-C5 antibodies, and C1 antibody 135/9. Moore & Sodroski [1996]
- 4A7C6: The relative affinity for denatured/native gp120 is 7.9 mutation 88 N/P impairs binding. Moore *et al.* [1994c]
- 4A7C6: C1 region epitope (88 N/P substitutions abrogates binding), but substitutions 380 G/F and 420 I/R also impaired binding. Moore *et al.* [1994d]
- 4A7C6: Bound preferentially to denatured IIIB gp120. Moore & Ho [1993]

**No.** 332 **MAb ID** 1D10

**HXB2 Location** gp160 (81–100) **Author Location** gp120 (81–100 LAI)

Keywords vaccine antigen design
• 133/192: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120,

References Pantophlet et al. 2004; Pantophlet et al.

2003b; Binley *et al.* 1998; Binley *et al.* 1997a; Trkola *et al.* 1996a; Moore & Sodroski 1996;

Moore et al. 1994d; Moore et al. 1994c;

Moore et al. 1993b; Niedrig et al. 1992b

**Ab Type** gp120 C1 **Research Contact** Matthias Niedrig

including 133/192. Pantophlet et al. [2004] (vaccine antigen design)

- 133/192: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)
- 133/192: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer. Binley et al.
- 133/192: Reciprocal binding inhibition with the antibody 4A7C6 - enhanced by some anti-C5 and-C1 antibodies. Moore & Sodroski [1996]
- 133/192: Does not neutralize JR-FL nor block gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]
- 133/192: The relative affinity for denatured/native gp120 is 1.8. Moore et al. [1994c]
- 133/192: C1 region substitutions 76P/Y, 113 D/A or R, 117 K/W, 420 I/R, 427 W/S impair binding, other substitutions enhanced binding. Moore et al. [1994d]
- 133/192: Epitope seems complex, binds multiple peptides weak neutralization of lab strain. Niedrig et al. [1992b]

No. 335 **MAb ID** 489.1(961) HXB2 Location gp160 (91-100)

Author Location gp120 (91–100 LAI)

**Epitope** ENFDMWKNDM

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

Research Contact C. Bruck, SKB, Belgium

References Moore et al. 1994c

- 489.1(961): NIH AIDS Research and Reference Reagent Pro-
- 489.1(961): The relative affinity for denatured/native gp120 is 1. Moore *et al*. [1994c]

**No.** 336

MAb ID 5B3

HXB2 Location gp160 (91–100)

Author Location gp120 (91-100 LAI)

**Epitope** ENFDMWKNDM

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp160

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

References Moore et al. 1994c; Beretta & Dalgleish 1994; Nakamura et al. 1992: Berman et al. 1991

- 5B3: The relative affinity of denatured/native gp120 is 8.3. Moore et al. [1994c]
- 5B3: Cross-blocks 1D10 in competitive IIIB-rsgp160 ELISA no neutralization - blocks IIIB-gp120 sCD4 binding - localized binding to residues 72-106. Nakamura et al. [1992]
- 5B3: Blocks gp120 -CD4 binding. Berman *et al.* [1991]

No. 337

MAb ID B10

HXB2 Location gp160 (91–100)

Author Location gp120 (91-100 LAI)

Epitope ENFDMWKNDM

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG1)

Ab Type gp120 C1

References Moore et al. 1994c; Abacioglu et al. 1994

- B10: There is FNM/FDM polymorphism in LAI-based peptides, and N is essential (J. P. Moore, per. comm.)
- B10: C1 region epitope boundaries mapped by peptide scanning, FNMW core. Abacioglu et al. [1994]
- B10: The relative affinity for denatured/native gp120 is 0.4. Moore et al. [1994c]

**No.** 338

MAb ID B2

HXB2 Location gp160 (91-100)

Author Location gp120 (91–100 LAI)

**Epitope** ENFDMWKNDM

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG2b)

Ab Type gp120 C1

References Binley et al. 1997a; Moore et al. 1994d; Moore et al. 1994c; Abacioglu et al. 1994;

Thali et al. 1993

- B2: There is FNM/FDM polymorphism in LAI-based peptides, and N is essential (J. P. Moore, per. comm.)
- B2: C1 region epitope boundaries mapped by peptide scanning, FNMW core. Abacioglu et al. [1994]
- B2: The relative affinity for denatured/native gp120 is 1.4. Moore *et al.* [1994c]

No. 339

MAb ID C6 (Ch6)

HXB2 Location gp160 (91–100)

Author Location gp120 (91–100 LAI)

**Epitope** ENFDMWKNDM

Subtype B

**Neutralizing** 

Immunogen vaccine

> Vector/Type: protein Strain: B clade LAI HIV component: gp160

Species (Isotype) mouse (IgG1)

Ab Type gp120 C1

References Pincus et al. 1996; Moore et al. 1994c; Aba- Species (Isotype) mouse (IgG) cioglu et al. 1994; Pincus & McClure 1993

- C6: There is FNM/FDM polymorphism in LAI-based peptides - N is essential (J. P. Moore, per. comm.)
- C6: NIH AIDS Research and Reference Reagent Program: 810.
- C6: Called Ch6 binds to gp120 but not to infected cells when linked to ricin A, the immunotoxin did not mediate cell killing – sCD4 has no effect. Pincus & McClure [1993]; Pincus et al. [1996]
- C6: C1 region epitope boundaries mapped by peptide scanning, FNMW core. Abacioglu et al. [1994]
- C6: The relative affinity for denatured/native gp120 is 0.9. Moore et al. [1994c]

**No.** 340

MAb ID MF49.1

HXB2 Location gp160 (91–100)

Author Location gp120 (91-100 LAI)

Epitope ENFDMWKNDM

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

• MF49.1: The relative affinity of denatured/native gp120 is 3.8. • T9: Binds to the C1 region – 45 W/S, 88 N/P, 256 S/Y, 262 Moore et al. [1994c]

**No.** 341

MAb ID T1.1

**HXB2 Location** gp160 (91–100)

**Author Location** gp120 (91–100 LAI)

**Epitope** ENFDMWKNDM

Subtype B

Neutralizing

Immunogen vaccine

*Vector/Type:* vaccinia HIV component:

gp160

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

Akerblom et al. 1990

- T1.1: C1 region the relative affinity for denatured/native gp120 is 1. Moore *et al.* [1994c]
- T1.1: Also reacted in solid phase with gp120(234-248) NGTG-PCTNVSTQCT. Akerblom et al. [1990]
- T1.1: No ADCC activity reactive peptide: NVTENFN-MWKNDMVEQ, IIIB. Broliden et al. [1990]

No. 342

MAb ID T7.1

HXB2 Location gp160 (91-100) Author Location gp120 (91-100 LAI)

Epitope ENFDMWKNDM

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Ab Type gp120 C1

References Moore et al. 1994d; Moore et al. 1994c; Bolm-

stedt et al. 1990; Akerblom et al. 1990

• T7.1: The relative affinity of denatured/native gp120 is 4.0. Moore et al. [1994c]

No. 343

MAb ID T9

HXB2 Location gp160 (91–100)

Author Location gp120 (91-100 LAI)

Epitope ENFDMWKNDM

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

Research Contact Lennart Akerblom, Britta Wahren and Jorma

Hinkula

References Binley et al. 1997a; Moore et al. 1994d;

Moore et al. 1994c; Bolmstedt et al. 1990; Akerblom et al. 1990

• T9: There are two HIV-Abs with the name T9, one binds to

- gp41, one to gp120.
- T9: The relative affinity of denatured/native gp120 is 7.9. Moore et al. [1994c]
- N/T, 475 M/S, 485 1.83, and 491 I/F enhanced binding, no substitution tested significantly inhibited. Moore et al. [1994d]

No. 344

MAb ID GV4D3

HXB2 Location gp160 (92–100)

Author Location gp120 (92–100 IIIB)

Epitope NFNMWKNDM

Neutralizing

Immunogen vaccine

Vector/Type: protein-Ab complex HIV com-

ponent: gp120-Mab complex

Species (Isotype) mouse

Ab Type gp120 C1

References Moore et al. 1994c; Broliden et al. 1990; Research Contact Patricia Earl and Christopher Broder, NIH References Denisova et al. 1996

> • GV4D3: When anti-V3 MAb M77 was bound to gp120 and used as an immunogen, it stimulated many MAbs to linear epitopes - MAbs GV4H4 and GV5F9 are homologous to GV4D3 and were generated in the same experiment. Denisova et al. [1996]

> > No. 345

MAb ID B27

**HXB2 Location** gp160 (93–96)

Author Location gp120 (94-97 BH10)

**Epitope** FNMW

Neutralizing no

gp160 Antibodies HIV Antibodies Tables

Immunogen vaccine No. 349 Vector/Type: protein Strain: B clade NL43 MAb ID D/5A11 HIV component: gp160 HXB2 Location gp160 (93–101) **Species (Isotype)** mouse (IgG1) Author Location gp120 (93-101 LAI) Epitope FNMWKNDMV Ab Type gp120 C1 Subtype B References Bristow et al. 1994; Abacioglu et al. 1994 • B27: C1 region – epitope boundaries mapped by peptide scan-Neutralizing no ning. Abacioglu et al. [1994] Immunogen vaccine • B27: C1 MAb generated in a study of the humoral immune re-Vector/Type: protein Strain: B clade LAI sponse to Baculovirus-expressed mis-folded rgp160 IIIB:NL43, HIV component: gp120 MicroGenSys. Bristow et al. [1994] Species (Isotype) mouse Ab Type gp120 C1 **No.** 346 References Bristow et al. 1994 MAb ID B9 • D/5A11: C1 MAb generated in a study of the humoral im-HXB2 Location gp160 (93-96) mune response to Baculovirus-expressed mis-folded rgp120 Author Location gp120 (93-96 LAI) and rgp160. Bristow et al. [1994] **Epitope** FNMW Subtype B **No.** 350 Neutralizing MAb ID D/6B2 Immunogen vaccine HXB2 Location gp160 (93-101) Vector/Type: protein Strain: B clade LAI Author Location gp120 (93–101 LAI) HIV component: gp160 **Epitope** FNMWKNDMV Species (Isotype) mouse (IgG1) Subtype B Ab Type gp120 C1 Neutralizing no References Abacioglu et al. 1994 Immunogen vaccine • B9: Binds C1 region – epitope boundaries mapped by peptide Vector/Type: protein Strain: B clade LAI scanning. Abacioglu et al. [1994] HIV component: gp120 Species (Isotype) mouse (IgG1) No. 347 Ab Type gp120 C1 MAb ID B35 References Bristow et al. 1994 **HXB2 Location** gp160 (93–98) • D/6B2: C1 MAb generated in a study of the humoral im-Author Location gp120 (94–99 BH10) mune response to Baculovirus-expressed mis-folded rgp120 Epitope FNMWKN and rgp160. Bristow et al. [1994] Neutralizing No. 351 Immunogen vaccine MAb ID B18 Vector/Type: protein Strain: B clade LAI HXB2 Location gp160 (101–110) HIV component: gp160 Species (Isotype) mouse (IgG1) Author Location gp120 (101–110 LAI) Ab Type gp120 C1 **Epitope** VEQMHEDIIS References Abacioglu et al. 1994 Subtype B • B35: C1 region – epitope boundaries mapped by peptide scan-Neutralizing ning. Abacioglu et al. [1994] Immunogen vaccine Vector/Type: protein Strain: B clade LAI **No.** 348 HIV component: gp160 MAb ID D/4B5 Species (Isotype) mouse (IgG2a) **HXB2 Location** gp160 (93–101) Ab Type gp120 C1 Author Location gp120 (93-101 LAI) References Moore et al. 1994c; Abacioglu et al. 1994 **Epitope** FNMWKNDMV • B18: C1 region – epitope boundaries mapped by peptide scan-Subtype B ning, HEDII core. Abacioglu et al. [1994] Neutralizing no • B18: The relative affinity for denatured/native gp120 is 1. Immunogen vaccine Moore et al. [1994c] Vector/Type: protein Strain: B clade LAI HIV component: gp120 No. 352 Species (Isotype) mouse MAb ID B20 Ab Type gp120 C1 HXB2 Location gp160 (101-110) References Bristow et al. 1994 Author Location gp120 (101-110 LAI) • D/4B5: C1 MAb generated in a study of the humoral im-**Epitope** VEQMHEDIIS mune response to Baculovirus-expressed mis-folded rgp120 Subtype B

> Neutralizing Immunogen vaccine

and rgp160. Bristow et al. [1994]

HIV component: gp160

Species (Isotype) mouse (IgG2a)

Ab Type gp120 C1

References Moore et al. 1994c; Abacioglu et al. 1994

- B20: C1 region epitope boundaries mapped by peptide scanning – HEDII core. Abacioglu et al. [1994]
- B20: The relative affinity for denatured/native gp120 is 1. Moore et al. [1994c]

**No.** 353

**MAb ID** MF39.1 (39.1)

HXB2 Location gp160 (101-110)

Author Location gp120 (101–110 LAI)

Epitope VEQMHEDIIS

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

References Moore et al. 1994c; Cook et al. 1994; Thiriart et al. 1989

- MF39.1: Called 39.1, and is probably the same as MF39.1 -MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon – MAbs against the N-terminal half of gp120 do not inhibit gp120 binding to GalCer - binding of GalCer to gp120 does not inhibit MAb binding. Cook et al. [1994]
- MF39.1: The relative affinity of denatured/native gp120 is 30. Moore et al. [1994c]

No. 354

**MAb ID** 187.2.1 (187.1)

HXB2 Location gp160 (101-120)

Author Location gp120 (101–120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

Research Contact Claudine Bruck and Clothilde Thiriart

References Moore et al. 1994d; Moore et al. 1994c; Cook et al. 1994; Moore & Ho 1993; Thiriart et al.

- 187.2.1: UK Medical Research Council AIDS reagent: ARP332.
- 187.2.1: Called 187.1, and is probably the same as 187.2.1 MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon - MAbs against the N-terminal half of gp120 do not inhibit gp120 binding to GalCer - binding of GalCer to gp120 does not inhibit MAb binding. Cook et al. [1994]
- 187.2.1: The relative affinity for denatured/native gp120 is 7 - mutations 113 D/A (not D/R) and 117 K/W impair binding. Research Contact Fulvia di Marzo Veronese Moore et al. [1994c]

Vector/Type: protein Strain: B clade LAI • 187.2.1: Called 187.1, and is probably the same as 187.2.1 – bound preferentially to denatured IIIB gp120. Moore & Ho [1993]

No. 355

**MAb ID** 37.1.1(ARP 327) (37.1)

HXB2 Location gp160 (101–120)

Author Location gp120 (101-120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

Research Contact Claudine Bruck

References Moore et al. 1994c; Moore & Ho 1993; Thiriart et al. 1989

- 37.1.1: UK Medical Research Council AIDS reagent: ARP327.
- 37.1.1: The relative affinity for denatured/native gp120 is 8.6 - mutations 113 D/R (not D/A) and 117 K/W impair binding. Moore et al. [1994c]
- 37.1.1: Called 37.1 bound preferentially to denatured IIIB gp120. Moore & Ho [1993]

No. 356

MAb ID 6D8

**HXB2 Location** gp160 (101–120)

Author Location gp120 (101-120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) rat

Ab Type gp120 C1

References Moore et al. 1994c; Nakamura et al. 1992;

Dowbenko et al. 1988

- 6D8: The relative affinity for denatured/native gp120 is 15 mutations 113 D/R and 113 D/A impair binding. Moore et al. [1994c]
- 6D8: Highly cross reactive with multiple stains by rgp120 ELISA. Nakamura et al. [1992]

No. 357

MAb ID M96

HXB2 Location gp160 (101-120)

Author Location gp120 (101–120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) rat (IgG2a)

Ab Type gp120 C1

References Moore et al. 1994d; Moore et al. 1994c;

di Marzo Veronese et al. 1992

gp160 Antibodies **HIV Antibodies Tables** 

• M96: C1 region – the relative affinity for denatured/native **Species (Isotype)** mouse (IgG) gp120 is 6. Moore *et al.* [1994c]

• M96: Immunoblot reactive for strains IIIB, 451, MN, RF, and RUTZ, di Marzo Veronese et al. [1992]

No. 358

**MAb ID** MF119.1

HXB2 Location gp160 (101-120) Author Location gp120 (101-120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG) Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

• MF119.1: The relative affinity for denatured/native gp120 is 30 - mutations 113 D/A, 113 D/R, and 117 K/W impair binding. Moore et al. [1994c]

**No.** 359

MAb ID MF4.1

HXB2 Location gp160 (101–120)

Author Location gp120 (101–120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG) Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

• MF4.1: The relative affinity for denatured/native gp120 is 8. Moore et al. [1994c]

**No.** 360

MAb ID MF53.1

HXB2 Location gp160 (101-120) Author Location gp120 (101-120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

**Neutralizing** 

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG) Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

• MF53.1: The relative affinity for denatured/native gp120 is 10. Moore et al. [1994c]

No. 361

MAb ID MF58.1

**HXB2 Location** gp160 (101–120)

Author Location gp120 (101–120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

No. 362

MAb ID MF77.1

HXB2 Location gp160 (101-120)

Author Location gp120 (101-120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

• MF77.1: The relative affinity for denatured/native gp120 is 11. Moore et al. [1994c]

No. 363

MAb ID T2.1

**HXB2 Location** gp160 (101–120)

Author Location gp120 (101–120 LAI)

Epitope VEQMHEDIISLWDQSLKPCV

Subtype B

**Neutralizing** 

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

Research Contact Lennart Akerblom, Britta Wahren and Jorma

References Moore et al. 1994d; Moore et al. 1994c; Bolm-

stedt et al. 1990: Akerblom et al. 1990

• T2.1: The relative affinity for denatured/native gp120 is .27 - mutations 113 D/R, 106 E/A, and 117 D/A impair binding. Moore *et al.* [1994c]

No. 364

**MAb ID** 11/65 (11/65a/5h)

HXB2 Location gp160 (102-121)

Author Location gp120 (311–321 HXB10)

Epitope EQMHEDIISLWDQSLKPCVK

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2b)

Ab Type gp120 C1

References Peet et al. 1998; McKeating et al. 1993b; McKeating et al. 1992a

- 11/65: UK Medical Research Council AIDS reagent: ARP3076.
- 11/65: Called 11/65a/5h The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic - these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind - 11/65 was not affected by V3 serine substitutions -

sponse relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]

• 11/65: Binds only soluble gp120, not virion bound – used to quantify gp120 shedding – (numbering is incorrect in original?) McKeating et al. [1992a]

No. 365

MAb ID W1

HXB2 Location gp160 (102-121) Author Location gp120 (102-121 LAI)

Epitope EQMHEDIISLWDQSLKPCVK

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

Research Contact D. Weiner, U. Penn.

References Moore et al. 1994c

• W1: The relative affinity for denatured/native gp120 is 6 mutations 113 D/A, 113 D/R, and 117 K/W impair binding. Moore et al. [1994c]

**No.** 366

MAb ID T11

HXB2 Location gp160 (102-125) Author Location gp120 (102-125)

Epitope EQMHEDIISLWDQSLKPCVKLTPL

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: oligomeric gp140

Species (Isotype) mouse

Ab Type gp120 C1

Research Contact R. Doms, Univ. of Pennsylvania

References Jagodzinski et al. 1996; Earl et al. 1994

- T11: The sulfated polysaccharide, curdlan sulfate (CRDS), binds to the Envelope of T-tropic viruses and neutralizes virus – deletion of the V3 loop from gp120 results in more potent T11 inhibition by CRDS. Jagodzinski et al. [1996]
- T11: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 367

MAb ID GV1A8

**HXB2 Location** gp160 (105–113) Author Location gp120 (105–113 IIIB)

Epitope HEDIISLWD

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein-Ab complex HIV component: gp120-Mab complex

Species (Isotype) mouse

Ab Type gp120 C1

References Denisova et al. 1996

mice injected with serine substituted gp120 had a reduced re- • GV1A8: When anti-V3 MAb M77 was bound to gp120 and used as an immunogen, it stimulated many MAbs to linear epitopes – MAbs GV7A4 and GV5H5 are homologous to GV1A8 and were generated in the same experiment. Denisova et al. [1996]

No. 368

MAb ID 11

HXB2 Location gp160 (111-120)

Author Location gp120 (101-120 LAI)

Epitope LWDQSLKPCV

Subtype B

**Neutralizing** 

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

• 11: The relative affinity for denatured/native gp120 is 7.8 – mutation 113 D/R impairs binding. Moore et al. [1994c]

No. 369

**MAb ID** 12G10

**HXB2 Location** gp160 (111–120)

Author Location gp120 (101–120 LAI)

Epitope LWDQSLKPCV

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

• 12G10: The relative affinity for denatured/native gp120 is 17 – mutation 117 K/W impairs binding. Moore et al. [1994c]

**No.** 370

**MAb ID** 135/9 (87-135/9)

HXB2 Location gp160 (111-120)

Author Location gp120 (111-120 LAI)

Epitope LWDQSLKPCV

Subtype B

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 C1

Research Contact Matthias Niedrig

References Yang et al. 2000; Kropelin et al. 1998; Binley et al. 1998; Binley et al. 1997a; Trkola et al. 1996a; Moore & Sodroski 1996; Moore et al. 1994d; Moore et al. 1994c; Niedrig et al.

1992h

• 135/9: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes - CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120

or gp140 – non-neutralizing MAbs C11, A32, 522-149, M90, Research Contact George Lewis and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 - MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang et al. [2000]

- 135/9: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer. Binley et al. [1998]
- 135/9: Noted to bind to C1 peptide HEDIISLWDQSLK blocks gp120 interaction with CD4+ cells - blocking activity is additive when combined with antibodies which bind in the C4 region of gp120 (F105, 388/389, and b12) Kropelin et al. [1998]
- 135/9: Binding is enhanced by some anti-C1 and anti-C5 antibodies - enhances binding of some anti-V3, anti-C4 and anti-V2 MAbs - 135/9 binds to predicted alpha-helix in C1. Moore & Sodroski [1996]
- 135/9: Does not neutralize JR-FL nor block gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]
- 135/9: The relative affinity for denatured/native gp120 is 15 - mutation 113 D/R impairs binding to native and denatured, 113 D/A only to denatured. Moore et al. [1994c]
- 135/9: Substitutions 106 E/A, 113 D/A or R, and 117 K/W impair binding, some substitutions enhance binding. Moore et al. [1994d]
- 135/9: Defines the epitope as gp120(114-123) MHEDIISLWD (core LWD?) - weak neutralization of lab strain. Niedrig et al. [1992b]

**No.** 371

MAb ID 7C10

HXB2 Location gp160 (111-120) Author Location gp120 (101-120 LAI)

**Epitope** LWDQSLKPCV

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

• 7C10: The relative affinity for denatured/native gp120 is 5.8 – mutation 117 K/W impairs binding. Moore et al. [1994c]

**No.** 372

MAb ID C4

HXB2 Location gp160 (111-120)

Author Location gp120 (101–120 LAI)

**Epitope** LWDQSLKPCV

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG1)

Ab Type gp120 C1

References Moore et al. 1994c: Moore & Ho 1993: Abacioglu et al. 1994

- C4: C1 region epitope boundaries mapped by peptide scanning, BH10 core IISLW. Abacioglu et al. [1994]
- C4: The relative affinity for denatured/native gp120 is 10. Moore *et al.* [1994c]
- C4: Bound preferentially to denatured IIIB gp120. Moore & Ho [1993]

**No.** 373

MAb ID MF46.1

HXB2 Location gp160 (111-120)

Author Location gp120 (101-120 LAI)

Epitope LWDQSLKPCV

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

References Moore et al. 1994c; Thiriart et al. 1989

• MF46.1: The relative affinity for denatured/native gp120 is 8.5. Moore et al. [1994c]

No. 374

MAb ID 6D5

**HXB2 Location** gp160 (122–141)

Author Location gp120 (122-141 LAI)

**Epitope** LTPLCVSLKCTDLKNDTNTN

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 V2

Research Contact S. Nigida and L. Arthur, NCI, Frederick, MD

References Moore et al. 1994d; Moore et al. 1994c

• 6D5: The relative affinity for denatured/native gp120 is 15 – mutations Delta119-205 and 125 L/G impair binding. Moore et al. [1994c]

No. 375

MAb ID B33

HXB2 Location gp160 (123–142)

Author Location gp120 (123-142 LAI)

**Epitope** TPLCVSLKCTDLGNATNTNS

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43

HIV component: gp160

**Species** (**Isotype**) mouse ( $IgG2b\kappa$ )

Ab Type gp120 V2

Research Contact Daniels

References Bristow et al. 1994; Abacioglu et al. 1994

• B33: UK Medical Research Council AIDS reagent: ARP304, gp160/41 binding.

• B33: There are two MAbs in the literature named B33, see also gp160(727-734) Abacioglu et al. [1994]

- B33: Epitope boundaries mapped by peptide scanning. Abacioglu et al. [1994]
- B27: C1 MAb generated in a study of the humoral immune response to Baculovirus-expressed mis-folded rgp160 IIIB:NL43, MicroGenSys. Bristow et al. [1994]

**No.** 376

MAb ID polyclonal (VEI1)

**HXB2 Location** gp160 (131–151) **Author Location** Env (131–151)

**Epitope** CTDLKNDTNTNSSSGRMMMEK

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Carlos et al. 1999

• Antibody response to the epitopes in a vaccine construct (VEI) containing peptides from 5 hypervariable regions of gp120 was detected in the sera of HIV-1 positive subjects, including sera from 6 non-subtype B infections – serum samples from San Francisco, Canada and Puerto Rico cohort showed presence of antibodies against all five VEI hypervariable regions, but most consistently against the V3 region peptide NNNTRKSIRIGP-GRAFYTTGDIGNIRQ. Carlos et al. [1999]

**No.** 377

MAb ID 35D10/D2

HXB2 Location gp160 (139-155)

Author Location gp120

Epitope NTKSSNWKEMDGEIK

Neutralizing L Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 V1

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002 **Keywords** antibody binding site definition and exposure, antibody generation, review, variant cross-

recognition or cross-neutralization

- 35D10/D2: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - several of the MAbs obtained were effective at neutralizing the autologous strain - C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were

highly type specific. He et al. [2002] (antibody binding site definition and exposure, antibody generation)

**No.** 378

MAb ID 40H2/C7

HXB2 Location gp160 (139–155)

Author Location gp120

Epitope NTKSSNWKEMDGEIK

Neutralizing L Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species** (**Isotype**) transgenic mouse ( $IgG2\kappa$ )

Ab Type gp120 V1

Research Contact Dr. Abraham Pinter, Public Health Research

Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002 Keywords antibody binding site definition and exposure,

antibody generation, review, variant crossrecognition or cross-neutralization

- 40H2/C7: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- 40H2/C7: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - several of the MAbs obtained were effective at neutralizing the autologous strain - C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were highly type specific. He et al. [2002] (antibody binding site definition and exposure, antibody generation)

No. 379

**MAb ID** 43A3/E4

HXB2 Location gp160 (139–155)

**Author Location** gp120

Epitope NTKSSNWKEMDGEIK

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 V1

• 35D10/D2: Transgenic mice (strain XenoMouse G2) carrying Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002

Keywords antibody binding site definition and exposure,

antibody generation, review, variant crossrecognition or cross-neutralization

- 43A3/E4: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- 43A3/E4: Transgenic mice (strain XenoMouse G2) carrying Research Contact Dr. Abraham Pinter, Public Health Research human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - several of the MAbs obtained were effective at neutralizing the autologous strain - C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were highly type specific. He et al. [2002] (antibody binding site definition and exposure, antibody generation)

**No.** 380

**MAb ID** 43C7/B9

HXB2 Location gp160 (139-155)

Author Location gp120

Epitope NTKSSNWKEMDGEIK

Neutralizing L Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 V1

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

> References Gorny & Zolla-Pazner 2004; He et al. 2002 Keywords antibody binding site definition and exposure, antibody generation, review, variant cross-

recognition or cross-neutralization

- 43C7/B9: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- 43C7/B9: Transgenic mice (strain XenoMouse G2) carrying **Research Contact** Dr. Abraham Pinter, Public Health Research human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - several of the MAbs obtained were effective at neutralizing the autologous strain - C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were highly type specific. He et al. [2002] (antibody binding site definition and exposure, antibody generation)

No. 381 MAb ID 45D1/B7 HXB2 Location gp160 (139-155) Author Location gp120

Epitope NTKSSNWKEMDGEIK

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species** (**Isotype**) transgenic mouse ( $IgG2\kappa$ )

Ab Type gp120 V1

Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002 **Keywords** antibody binding site definition and exposure,

antibody generation, review, variant crossrecognition or cross-neutralization

- 45D1/B7: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- 45D1/B7: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - several of the MAbs obtained were effective at neutralizing the autologous strain - C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were highly type specific. He et al. [2002] (antibody binding site definition and exposure, antibody generation)

No. 382

**MAb ID** 46E3/E6

HXB2 Location gp160 (139–155)

Author Location gp120

Epitope NTKSSNWKEMDGEIK

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species** (**Isotype**) transgenic mouse ( $IgG2\kappa$ )

Ab Type gp120 V1

Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002 **Keywords** antibody binding site definition and exposure, antibody generation, review, variant crossrecognition or cross-neutralization

• 46E3/E6: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)

• 46E3/E6: Transgenic mice (strain XenoMouse G2) carrying **Research Contact** Dr. Abraham Pinter, Public Health Research human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - several of the MAbs obtained were effective at neutralizing the autologous strain - C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were highly type specific. He et al. [2002] (antibody binding site definition and exposure, antibody generation)

**No.** 383 **MAb ID** 58E1/B3

HXB2 Location gp160 (139-155)

Author Location gp120

Epitope NTKSSNWKEMDGEIK

Neutralizing L Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 V1

Research Contact Dr. Abraham Pinter, Public Health Research

Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002 Keywords antibody binding site definition and exposure, antibody generation, review, variant crossrecognition or cross-neutralization

- 58E1/B3: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - several of the MAbs obtained were effective at neutralizing the autologous strain - C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were highly type specific. He et al. [2002] (antibody binding site definition and exposure, antibody generation)

**No.** 384

MAb ID 64B9/A6

HXB2 Location gp160 (139-155)

Author Location gp120

Epitope NTKSSNWKEMDGEIK

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 V1

Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002 **Keywords** antibody binding site definition and exposure, antibody generation, review, variant crossrecognition or cross-neutralization

- 64B9/A6: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- 64B9/A6: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - several of the MAbs obtained were effective at neutralizing the autologous strain - C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were highly type specific. He et al. [2002] (antibody binding site definition and exposure, antibody generation)

No. 385

MAb ID 69D2/A1

**HXB2 Location** gp160 (139–155)

Author Location gp120

Epitope NTKSSNWKEMDGEIK

Neutralizing L Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 V1

58E1/B3: Transgenic mice (strain XenoMouse G2) carrying Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002

**Keywords** antibody binding site definition and exposure, antibody generation, review, variant crossrecognition or cross-neutralization

- 69D2/A1: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- 69D2/A1: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - several of the MAbs obtained were effective at neutralizing the autologous strain - C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were

highly type specific. He *et al.* [2002] (antibody binding site definition and exposure, antibody generation)

**No.** 386 **MAb ID** 82D3/C3

HXB2 Location gp160 (139–155)

**Author Location** gp120

**Epitope NTKSSNWKEMDGEIK** 

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species** (**Isotype**) transgenic mouse ( $IgG2\kappa$ )

Ab Type gp120 V1

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

**References** Gorny & Zolla-Pazner 2004; He *et al.* 2002 **Keywords** antibody binding site definition and exposure, antibody generation, review, variant cross-recognition or cross-neutralization

- 82D3/C3: In an extensive review of gp120 binding MAbs in this database, V1-specific MAbs are noted to be unusual, but in one study transgenic mouse immunization with SF162 yeilded ten V1 MAbs with highly strain-specific autologous NAb responses, and V1 was immunodominant. Human MAbs to V1 may be difficult to isolate due to selection with heterologous strains. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization, review)
- 82D3/C3: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 several of the MAbs obtained were effective at neutralizing the autologous strain C-term-V1- region was immunodominant in these mice and the ten V1-specific MAbs could potently neutralize autologous strain SF162 but were highly type specific. He *et al.* [2002] (antibody binding site definition and exposure, antibody generation)

**No.** 387

MAb ID 2H1B

HXB2 Location gp160 (155–161)

Author Location gp120 (370–376 HIV2ROD)

Epitope RNISFKA

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: HIV-2 ROD

Species (Isotype) mouse

Ab Type gp120 C3

References Matsushita et al. 1995

2H1B: Binds in WB, but binds poorly to Env on the cell surface.
 Matsushita et al. [1995]

**No.** 388

**MAb ID** 697-D (697D, 697-30D)

HXB2 Location gp160 (161–180)

Author Location gp120 (161–180 IIIB)

Epitope ISTSIRGKVQKEYAFFYKLD

Neutralizing P (weak)

**Immunogen** HIV-1 infection **Species** (**Isotype**) human ( $IgG1\lambda$ )

Ab Type gp120 V2

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

las01@mcrcr6.med.nyu) (NYU Med. Center) or Cellular Products Inc, Buffalo NY

References Selvarajah et al. 2005; Gorny & Zolla-Pazner

2004; He et al. 2002; Maksiutov et al. 2002; Edwards et al. 2002; Nyambi et al. 2000; Hioe et al. 2000; Gorny et al. 2000; Stamatatos & Cheng-Mayer 1998; Nyambi et al. 1998; Parren et al. 1997b; Fouts et al. 1997; Binley et al. 1997a; Trkola et al. 1996a; Moore & Ho 1995; Forthal et al. 1995; Gorny et al. 1994

**Keywords** ADCC, antibody binding site definition and exposure, co-receptor, enhancing activity, review, subtype comparisons, vaccine anti-

gen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-

neutralization

- 697-D: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. V2 MAb 697-D did not bind to mCHO and had diminished binding to GDMR, while V2 MAb 8.22.2 bound to GDMR but not mCHO. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)
- 697-D: In a review of Envelope binding MAbs in this database, V2-specific MAbs are noted to have some ability to neutralize HIV-1, but generally weak with limited cross-reactivity; it weakly neutralizes some primary but not TCLA strains. 697-D is the best characterized of the anti-V2 MAbs, and binds weakly and sporadically to isolates from clades A-D. Gorny & Zolla-Pazner [2004] (variant cross-recognition or crossneutralization, review, subtype comparisons)
- 697-D: Called 697D Truncation of the gp41 cytoplasmic domain of X4, R5, and X4R5 viruses forces a conformation that more closely resembles the CD4 bound state of the external Envelope, enhancing binding of CD4i MAbs 17b and 48d and of CD4BS MAbs F105, b12, and in most cases of glycosylation site dependent MAb 2G12 and the anti-gp41 MAb 246D in contrast, binding of the anti-V2 MAb 697D and the anti-V3 MAb 694/98D were not affected viruses bearing the truncation were more sensitive to neutralization by MAbs 48d, b12, and 2G12 the anti-C5 MAb 1331A was used to track levels of cell surface expression of the mutated proteins. Edwards *et al.* [2002] (antibody binding site definition and exposure)
- 697-D: Called 697D Transgenic mice carrying human genes allowing production of fully human MAbs were used to rapidly create a panel of anti-HIV gp120 MAb producing hybridomas by immunization with HIV SF162 gp120 – the previously described human MAbs 5145A, 4117C and 697D were used as controls. He *et al.* [2002]

• 697-D: This epitope is similar to a fragment of the human protein macrophage colony stimulating factor I receptor SISIR-LKVOK. Maksiutov et al. [2002]

- 697-D: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared – no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer - V2 MAbs 697-D, 1357 and 1361 favored the monomer by approximately 2 fold. Gorny et al. [2000] (antibody binding site definition and exposure)
- 697-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses - CD4BS MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells – V2 MAb 697-D did not effect proliferation. Hioe et al. [2000]
- 697-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 anti-V2 MAbs, which showed weak and sporadic binding, with the most frequent binding to C and D clades. Nyambi et al. [2000] (subtype comparisons)
- 697-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H - V2 Abs 697-D, 1361, and 1357 tended to bind weakly with a similar pattern of specificity to virions, and bound well to soluble gp120: weak binding to 1/4 B clade viruses (CA5), and weak binding to viruses from subtype A and D. Nyambi et al. [1998] (subtype comparisons)
- 697-D: Called 697-30D deleting the V2 loop of neutralizationresistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F - deletion of V1 or V2 did not enable neutralization by V2 MAbs G3.4, G3.136, or 687-30D. Stamatatos & Cheng-Mayer [1998] (variant cross-recognition or crossneutralization)
- 697-D: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding - 697-D bound monomer, did not bind oligomer or neutralize JRFL. Fouts et al. [1997] (antibody binding site definition and exposure)
- 697-D: Does not neutralize TCLA strains but neutralizes some primary isolates weakly. Parren et al. [1997b] (variant crossrecognition or cross-neutralization)
- 697-D: Partial inhibition of gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]
- 697-D: Not neutralizing, no ADCC activity, and no viral enhancing activity. Forthal et al. [1995] (ADCC, enhancing activity)
- 697-D: Review: called 697/30D neutralizes some primary, but not lab adapted strains. Moore & Ho [1995] (variant crossrecognition or cross-neutralization, review)
- 697-D: Conformational with weak reactivity to V2 peptide IST-SIRGKVQKEYAFFYKLD - neutralized 3/4 primary isolates, but none of 4 lab strains - V2 substitutions 176/177 FY/AT, 179/180 LD/DL, 183/184 PI/SG, and 192-194 YSL/GSS abrogate binding - anti-C4 MAbs G3-536 and G45-60 enhance

binding - mild oxidation of carbohydrate moieties inhibits binding. Gorny et al. [1994] (antibody binding site definition and exposure)

No. 389

MAb ID 6C4/S

HXB2 Location gp160 (162–169)

Author Location gp120 (BH10)

**Epitope** STSIRGKV

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype)

Research Contact S. Ranjbar (NIBSC, UK)

References Moore et al. 1993a

• 6C4/S: UK Medical Research Council AIDS reagent: ARP3049.

**No.** 390

MAb ID C108G (C108g)

**HXB2 Location** gp160 (162–169)

Author Location gp120 (162–169 HXB2)

**Epitope** STSIRGKV

Subtype B

Neutralizing L

Immunogen HIV-1 infection

**Species** (**Isotype**) chimpanzee ( $IgG1\kappa$ )

Ab Type gp120 V2

**Research Contact** S. Tilley, Public Health Research Institute,

NY, NY References Pinter et al. 2005: Gorny & Zolla-Pazner

> 2004; Alsmadi & Tilley 1998; Mondor et al. 1998; Ugolini et al. 1997; Warrier et al. 1996; Warrier et al. 1995; Wu et al. 1995; Warrier

et al. 1994

Keywords ADCC, antibody binding site definition and exposure, antibody generation, antibody interactions, antibody sequence, variable domain, review, variant cross-recognition or

cross-neutralization

• C108G: This MAb is type-specific and neutralizes BaL and HXB2. It is the most potent anti-V2 MAb, is glycan dependent, and contrary to earlier reports requires disulfide bonds. Neutralization by C108g is not mediated by CD4 or CCR5 receptor blockage on the cell surface. Binding to CD4 was inhibited by b12, but not by C108g. Binding to CCR5 was completely inhibited by two V3 MAbs, 4117C and 2219, and was substantially inihibited by 2G12, but was not inhibited by C108g.JR-FL is a neutralization resistant strain; modification of JRFL at positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The MAb 10/76b, that binds to a linear V2 epitope that is unaffected by deglycosylation or reduction eliminating disulfinde bonds, could only weakly neutralize this modified JR-FL. Similarly SF162 substitutions in the neutralization sensitive virus SF162 GVK->NMK (167-169) plus the glycoslyation site at 160, created a G108g neutralization sensitive virus. In contrast, 10/76b binds to the NMK substituted variant, but addition of the glycosylation site inhibited binding. Pinter et al. [2005] (antibody binding

site definition and exposure, variant cross-recognition or • 10/76b: This study is about the MAb C108g, and 10/76b was a cross-neutralization)

- C108G: This MAb is unusual among V2-directed MAbs. It is glycan dependent and can neutralize both a primary isolate (BaL and a TCLA (IIIB) strain. Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, review)
- C108G: A study of 6 anti-Env MAbs and their ability to bind or direct ADCC against target cells infected with IIIB, MN, SF-2, and RF – C108G bound and directed lysis against only IIIB – this is first demonstration of ADCC directed by a V2 specific MAb. Alsmadi & Tilley [1998] (ADCC, variant crossrecognition or cross-neutralization)
- C108G: Inhibits HX10 binding to both CD4 positive and negative HeLa cells. Mondor et al. [1998]
- C108G: Viral binding inhibition by C108G was correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5) Ugolini et al. [1997] (antibody binding site definition and exposure)
- C108G: Synergistic neutralization of HIV-1 when combined with anti-V3 MAbs 0.5beta and C311E, or anti-CD4BS MAbs, 1125H and 5145A - neutralization further enhanced by presence of both 1125H and 0.5beta. Warrier et al. [1996] (antibody interactions)
- C108G: Characterization of MAb variable region. Warrier et al. [1995] (antibody sequence, variable domain)
- C108G: Strain specificity: LAI, BaL, HXB2 conformational character - glycosylation site at 160 critical - mutation of conserved glycosylation site at 156 increased epitope exposure. Wu et al. [1995] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- C108G: Chimps were infected with HIV-1 IIIB, and this high affinity MAb was obtained from an Epstein-Barr virus transformed B-cell line. It gave potent neutralization of HIV-1 IIIB. Binding was not affected by reduction of disulfide bonds. Binding was disrupted by removal of N-linked glycans. The peptide binds with lower affinity than glycosylated Env. Warrier et al. [1994] (antibody binding site definition and exposure, antibody generation)

No. 391

MAb ID 10/76b

HXB2 Location gp160 (162–170)

**Author Location** gp120 (162–171 BH10)

Epitope STSIRGKVQ Neutralizing L (HXB10)

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2a)

Research Contact Jane McKeating

References Pinter et al. 2005; McKeating et al. 1996; Wu et al. 1995; Shotton et al. 1995; McKeating et al. 1993a; McKeating et al. 1993b

**Keywords** antibody binding site definition and exposure, antibody generation, variant cross-recognition

or cross-neutralization

• 10/76b: UK Medical Research Council AIDS reagent: ARP3077.

- control. C108g is type-specific and neutralizes BaL and HXB2. It is the most potent anti-V2 MAb, and is glycan dependent and contrary to earlier reports requires disulfide bonds. Neutralization by C108g is not mediated by CD4 or CCR5 receptor blockage on the cell surface. JR-FL is a neutralization resistant strain; modification of JRFL at positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The MAb 10/76b, that binds to a linear V2 epitope that is unaffected by deglycosylation or reduction eliminating disulfinde bonds, could only weakly neutralize this modified JR-FL. Similarly SF162 substitutions in the neutralization sensitive virus SF162 GVK->NMK (167-169) plus the glycoslyation site at 160, created a G108g neutralization sensitive virus. In contrast, 10/76b binds to the NMK substituted variant, but addition of the glycosylation site inhibited binding. Pinter et al. [2005] (antibody binding site definition and exposure)
- 10/76b: Neutralizes HXB2, but fails to neutralize chimeric virus with gp120 from primary isolates in an HXB2 background. McKeating et al. [1996]
- 10/76b: Cross-competes with MAbs 10/76b and 11/4b HXB2 neutralization escape mutant has the substitution I/T at residue 165. Shotton et al. [1995] (antibody binding site definition and exposure)
- 10/76b: Included in cross-competition and neutralization studies. Shotton et al. [1995] (antibody binding site definition and exposure)
- 10/76b: HX10 strain specificity binds native, deglycosylated, or denatured gp120. Wu et al. [1995] (antibody binding site definition and exposure, variant cross-recognition or crossneutralization)
- 10/76b: This MAb was obtained from a hybridoma cell line. An R to L substitution abrogated binding. Human sera recognize the 10/76 epitope. McKeating et al. [1993b] (antibody generation)

No. 392

MAb ID 11/41e

HXB2 Location gp160 (162–170)

**Author Location** gp120 (162–171)

Epitope STSIRGKVQ

Neutralizing L (HXB10)

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

**Species (Isotype)** rat (IgG1)

References Wu et al. 1995; Shotton et al. 1995; McKeating et al. 1993b

- 11/41e: Included in cross-competition and neutralization studies. Shotton et al. [1995]
- 11/41e: HX10 strain specificity binds native and deglycosylated gp120. Wu *et al.* [1995]
- 11/41e: R to L abrogated binding human sera recognize the epitope. McKeating et al. [1993b]

**No.** 393

**MAb ID** 11/4b

HXB2 Location gp160 (162–170)

**Author Location** gp120 (162–171)

**Epitope** STSIRGKVQ **Neutralizing** L (HXB10) **Immunogen** vaccine

Vector/Type: protein Strain: B clade BH10 HIV component: gp120

Species (Isotype) rat (IgG2a)

**References** Moore & Sodroski 1996; Wu *et al.* 1995; Shotton *et al.* 1995; McKeating *et al.* 1993b

- 11/4b: Linear V2 epitope reciprocal binding enhancement of anti-V2 discontinuous epitope antibodies (in contrast to BAT085) and CD4 inducible antibody 48d. Reciprocal inhibits BAT085 binding inhibits CRA-3 binding CRA-3 does not inhibit 11/4b. Moore & Sodroski [1996]
- 11/4b: Cross-competes with MAbs 10/76b and 11/4c HXB2 neutralization escape mutant has the substitution I/T at residue 165. Shotton *et al.* [1995]
- 11/4b: HXB10 strain specificity binds native, deglycosylated, or denatured gp120. Wu et al. [1995]
- 11/4b: A change from R to L abrogated binding human sera recognize epitope. McKeating *et al.* [1993b]

**No.** 394

MAb ID RSD-33

HXB2 Location gp160 (162–170)

Author Location gp120 (162-171)

Epitope STSIRGKVQ

Neutralizing

Immunogen vaccine

*Vector/Type:* protein *Strain:* B clade BH10 *HIV component:* gp120

Species (Isotype)

**Research Contact** R. Daniels (NIMR, UK) **References** Moore *et al.* 1993a

No. 395

**MAb ID** 11/4c (11/4c/1j/4j) **HXB2 Location** gp160 (162–170)

**Author Location** gp120 (152–181)

Epitope STSIRGKVQ

Neutralizing L (HXB2)

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10 HIV component: gp120

Species (Isotype) rat (IgG2a)

Ab Type gp120 V2

References Peet et al. 1998; Shotton et al. 1995; Wu et al. 1995; McKeating et al. 1993b

- 11/4c: UK Medical Research Council AIDS reagent: ARP3035.
- 11/4c: Called 11/4c/1j/4j The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic – these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind – 11/4c was not affected by V3 serine substitutions – mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]
- 11/4c: Cross-competes with MAbs 10/76b and 11/4b HXB2 neutralization escape mutant has the substitution I/T at residue 165. Shotton *et al.* [1995]

 11/4c: HX10 strain specificity – binds native, deglycosylated, or denatured gp120. Wu et al. [1995]

 11/4c: R to L substitution abrogated binding – human sera recognize epitope. McKeating et al. [1993b]

**No.** 396

**MAb ID** 8.22.2

HXB2 Location gp160 (162–178)

**Author Location** gp120

Epitope TTSIRDKVQKEYALFYK

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species** (**Isotype**) transgenic mouse ( $IgG2\kappa$ )

Ab Type gp120 V2

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

**References** Selvarajah *et al.* 2005; Pinter *et al.* 2004; Pantophlet *et al.* 2004; Gorny & Zolla-Pazner

tophlet *et al.* 2004; Gorny & Zolla-Pazner 2004; Maksiutov *et al.* 2002; He *et al.* 2002

**Keywords** antibody binding site definition and exposure, antibody generation, review, subtype comparisons, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-neutralization

- 8.22.2: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. V2 MAb 697-D did not bind to mCHO and had diminished binding to GDMR, while V2 MAb 8.22.2 bound to GDMR but not mCHO. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)
- 8.22.2: This broad review of anti-Envelope MAbs notes that V2 MAbs are generally weakly neutralizing at best, and somewhat strain specific. 8.22.2 weakly neutralizes SF162. Gorny & Zolla-Pazner [2004] (review)
- 8.22.2: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including 8.22.2. Pantophlet *et al.* [2004] (vaccine antigen design)
- 8.22.2: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and

2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Three anti-V2 MAb were tested – 8.22.2 weakly neutralized SF162, and did not neutralize JRFL at all. Pinter *et al.* [2004] (variant cross-recognition or cross-neutralization)

- 8.22.2 : Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 several of the MAbs obtained were effective at neutralizing the autologous strain 8.22.2 was the only V2-specific MAb created and it could cross-compete with MAb 697D 8.22.2 could cross-react with BaL and JR-FL, two B clade R5 strains, but not B clade X4 or E clade viruses, and it could weakly neutralize autologous strain SF162. He et al. [2002] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)
- 8.22.2: This epitope is similar to a fragment of the human protein macrophage colony stimulating factor I receptor SISIR-LKVQK. Maksiutov et al. [2002]

No. 397

MAb ID 12b

HXB2 Location gp160 (162-181)

Author Location gp120 (162-181)

Epitope STSIRGKVQKEYAFFYKLDI

**Neutralizing** L (HXB10) **Immunogen** vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2a)

Ab Type gp120 V2

References Maksiutov et al. 2002; McKeating et al. 1996;

Shotton et al. 1995

- 12b: This epitope is similar to a fragment of the human protein macrophage colony stimulating factor I receptor SISIR-LKVQK. Maksiutov et al. [2002]
- 12b: Neutralizes HXB2, but fails to neutralize chimeric virus with gp120 from primary isolates in an HXB2 background. McKeating et al. [1996]
- 12b: V2 MAb neutralized HXB2 position 179-180 LD to DL abrogates binding competes with 60b, but not 74. Shotton *et al.* [1995]

No. 398

**MAb ID** G3-136 (G3.136)

**HXB2 Location** gp160 (170–180)

Author Location gp120 (170–180 IIIB)

Epitope QKEYAFFYKLD

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) mouse (IgG)

Ab Type gp120 V2

Research Contact Tanox Biosystems Inc and David Ho,

ADARC, NY

References Pantophlet et al. 2003b; Zwick et al. 2003; Ly & Stamatatos 2000; Stamatatos & Cheng-Mayer 1998; Parren et al. 1998a; Wyatt et al. 1997; Ditzel et al. 1997; Stamatatos et al. 1997; Binley et al. 1997a; Poignard et al. 1996a; Moore & Sodroski 1996; Stamatatos & Cheng-Mayer 1995; Sattentau & Moore 1995; Yoshiyama et al. 1994; Moore et al. 1993a; Moore & Ho 1993; Thali et al. 1993; Pirofski

Keywords antibody interactions, vaccine antigen design

G3-136: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of non-neutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)

et al. 1993; Fung et al. 1992

- G3-136: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the V2 MAb used. Zwick *et al.* [2003] (antibody interactions)
- G3-136: Called G3.136 SF162 is a neutralization-resistant HIV-1 isolate N-linked glycosylation modifications in the V2 loop of the SF162 gp120 revealed that these sites prevent neutralization by CD4BS MAbs (IgG1b12 and IgGCD4), and protect against neutralization by V3 MAbs (447-D and 391-95D) V2-region glycosylation site mutations did not alter neutralization resistance to V2 MAbs (G3.4 and G3.136) or CD4i MAbs (17b and 48d) V2 glycosylation site modification allows infection of macrophages, probably due to glycosylated forms requiring fewer CCR5 molecules for viral entry. Ly & Stamatatos [2000]
- G3-136: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- G3-136: Called G3.136 deleting the V2 loop of neutralization-resistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F deletion of V1 or V2 did not enable neutralization by V2 MAbs G3.4, G3.136, or 687-30D. Stamatatos & Cheng-Mayer [1998]
- G3-136: Called G3.136 does not mediate gp120 virion dissociation in contrast to anti-V2 MAb G3-4 not neutralizing for SF162 or SF128A in either primary macrophages or PBMC. Stamatatos *et al.* [1997]

• G3-136: Binds both gp120 and soluble gp120+gp41 complex efficiently, suggesting its gp120 epitope is not blocked by gp41 binding. Wyatt et al. [1997]

- G3-136: Described epitope as STSIRGKVKEYAFFYKLDI - binds oligomer - binding of V2 MAbs G3-136, G3-4 or BAT123 did not significantly alter gp120 dissociation from virus or expose the gp41 epitope of MAb 50-69, in contrast to anti-V3 MAbs. Poignard et al. [1996a]
- G3-136: Bound preferentially to the monomeric rather than oligomeric form of LAI gp120 - neutralizes cell free Hx10. Sattentau & Moore [1995]
- G3-136: The binding of conformation-dependent anti-V2, anti-V3, and anti-CD4BS MAbs to monomeric and virionassociated gp120 from HIV-1 isolates with differences in cell tropism was studied - V2 loop epitopes are somewhat occluded in the oligomeric gp120 epitopes on the virion surface relative to the gp120 monomer as indicated by an increase in the halfmaximal binding values to macrophage-tropic isolates SF162 and SF128a - anti-V2 MAbs G3-4 and G3.136 don't bind to T-cell tropic SF2. Stamatatos & Cheng-Mayer [1995]
- G3-136: HIV-1 RF V2 substitutions 177 Y/H and 179 L/P in the V2 loop of RF reduce affinity. Yoshiyama et al. [1994]
- G3-136: Conformational, does not bind well to denatured gp120 - not reactive with SF-2 gp120, and does not inhibit HIV-1 sera from binding to IIIB gp120. Moore & Ho [1993]
- G3-136: Marginal binding to peptide, binding inhibited by 183/184 PI/SG substitution. Moore et al. [1993a]
- G3-136: Binding enhanced by selected antibodies to C1, C4, C5, V3 and anti-CD4 binding site MAbs – enhances binding of selected V3, C4 and anti-CD4 binding site MAbs. Moore et al. [1993a]
- G3-136: V2 region binds and neutralizes IIIB and RF in CEM-SS cells, but not MN - neutralization activity against a few primary isolates in PBMC - sCD4 binding inhibits binding (contrast with BAT085) - deglycosylation or reduction of gp120 by DTT diminishes reactivity. Fung et al. [1992]

**No.** 399

**MAb ID** G3-4 (G3.4)

HXB2 Location gp160 (170–180)

**Author Location** gp120 (170–180 BH10)

Epitope QKEYAFFYKLD

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

**Species** (**Isotype**) mouse ( $IgG2b\kappa$ )

Ab Type gp120 V2

Research Contact Tanox Biosystems Inc and David Ho,

ADARC, NY

References Gorny et al. 2005; Pantophlet et al. 2004; Mc-Caffrey et al. 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Srivastava et al. 2002; Ly & Stamatatos 2000; Stamatatos & Cheng-Mayer 1998; Parren et al. 1998a; Wyatt et al. 1997; Ditzel et al. 1997; Stamatatos et al. 1997; Binley et al. 1997a; Poignard et al. 1996a; Moore & Sodroski 1996; Jagodzinski et al. 1996; Sattentau & Moore 1995; Wu et al. 1995; Stamatatos & Cheng-Mayer 1995; Yoshiyama

et al. 1994; Thali et al. 1994; Gorny et al. 1994: Moore et al. 1994b: Moore et al. 1993a: Thali et al. 1993; Sattentau et al. 1993; Sullivan et al. 1993; Moore & Ho 1993; McKeating et al. 1992a; Fung et al. 1992; Ho et al. 1992; Ho et al. 1991a

**Keywords** antibody binding site definition and exposure, antibody interactions, vaccine antigen design

- G3-4: 2909 is a human anti-Env NAb that was selected by neutralization assay and binds to the quaternary structure on the intact virion. G3-4 was used as a positive control for defining the binding properties of 2909. Gorny et al. [2005]
- G3-4: Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The SF2 and all five glycan mutants were resistant to G3-4. V3 glycans tended to shield V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans shielded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004] (antibody binding site definition and exposure, vaccine antigen design)
- G3-4: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including G3-4. Pantophlet *et al.* [2004] (vaccine antigen design)
- G3-4: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional Nlinked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)
- G3-4: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the V2 MAbs used. Zwick et al. [2003] (antibody interactions)
- G3-4: Called G3.4 Oligomeric gp140 (o-gp140) derived from R5 primary isolate US4 was characterized for use as a vaccine reagent - antigen capture ELISA was used to compare the antigenicity of gp120 and o-gp140 using a panel of well

et al. [2002]

- G3-4: Called G3.4 SF162 is a neutralization-resistant HIV-1 isolate - N-linked glycosylation modifications in the V2 loop of the SF162 gp120 revealed that these sites prevent neutralization by CD4BS MAbs (IgG1b12 and IgGCD4), and protect against neutralization by V3 MAbs (447-D and 391-95D) – V2region glycosylation site mutations did not alter neutralization resistance to V2 MAbs (G3.4 and G3.136) or CD4i MAbs (17b and 48d) – V2 glycosylation site modification allows infection of macrophages, probably due to glycosylated forms requiring fewer CCR5 molecules for viral entry. Ly & Stamatatos [2000]
- G3-4: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated – authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- G3-4: Called G3.4 Deleting the V2 loop of neutralizationresistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F - deletion of V1 or V2 did not enable neutralization by V2 MAbs G3.4, G3.136, or 687-30D. Stamatatos & Cheng-Mayer [1998]
- G3-4: Called G3.4 mediates gp120 virion dissociation in contrast to anti-V2 MAb G3-136 - not neutralizing for SF162 or SF128A in either primary macrophages or PBMC. Stamatatos et al. [1997]
- G3-4: Binds both gp120 and soluble gp120+gp41 complex efficiently, suggesting its gp120 epitope is not blocked by gp41 binding. Wyatt et al. [1997]
- G3-4: The sulfated polysaccharide curdlan sulfate (CRDS) binds to the Envelope of T-tropic viruses and neutralizes virus - deletion of the V3 loop from gp120 results in more potent G3-4 binding inhibition by CRDS – G3-4 epitope described as 176-184 FYKLDIIPI and 191-193 YSL. Jagodzinski et al. [1996]
- G3-4: Binding enhanced by selected antibodies to C1, C4, C5, V3 and anti-CD4 binding site MAbs - enhances binding of selected V3, C4 and anti-CD4 binding site MAbs. Moore & Sodroski [1996]
- G3-4: Described epitope as STSIRGKVKEYAFFYKLDI binds oligomer - binding of V2 MAbs G3-136, G3-4 or BAT085 did not significantly alter gp120 dissociation from virus or expose the gp41 epitope of MAb 50-69, in contrast to anti-V3 MAbs. Poignard et al. [1996a]
- G3-4: Bound preferentially to the monomeric rather than oligomeric form of LAI gp120 - neutralizes Hx10 cell-free virus. Sattentau & Moore [1995]
- G3-4: The binding of conformation-dependent anti-V2, anti-V3, and anti-CD4BS MAbs to monomeric and virionassociated gp120 from HIV-1 isolates with differences in cell tropism was studied - V2 loop epitopes are somewhat occluded in the oligomeric gp120 epitopes on the virion surface relative to the gp120 monomer as indicated by an increase in the halfmaximal binding values to macrophage-tropic isolates SF162 and SF128a - anti-V2 MAbs G3-4 and G3.136 don't bind to T-cell tropic SF2. Stamatatos & Cheng-Mayer [1995]

- characterized MAbs G3.4 recognized o-gp140. Srivastava G3-4: Reactive with BH10, RF, and MN binds native, but not denatured or deglycosylated gp120, binds to deglycosylated V1V2 fusion protein, suggesting importance of glycans outside the V1V2 region. Wu et al. [1995]
  - G3-4: Weakly neutralizing, IC 50 = 53 mug/ml. Gorny et al. [1994]
  - G3-4: Conformationally sensitive sporadic cross-reactivity among, and outside, B clade gp120s. Moore et al. [1994b]
  - G3-4: gp41 mutation (582 A/T) that reduces neutralization of anti-CD4 binding site MAbs does not alter G3-4s ability to neutralize. Thali et al. [1994]
  - G3-4: Neutralizes RF substitutions 177 Y/H and 179 L/P in the V2 loop of RF reduce affinity and result in neutralization escape. Yoshiyama et al. [1994]
  - G3-4: Conformational, does not bind well to denatured gp120 - not reactive with SF-2 gp120, and does not inhibit HIV-1 sera from binding to IIIB gp120. Moore & Ho [1993]
  - G3-4: V2 region, marginal binding to peptide, binding inhibited by 183/184 PI/SG substitution. Moore et al. [1993a]
  - G3-4: Increased binding in the presence of sCD4. Sattentau et al. [1993]
  - G3-4: Substitutions in residues 176 to 184 affect MAb recognition – substitutions in V2 can result in gp120-gp41 dissociation. Sullivan et al. [1993]
  - G3-4: Neutralizes IIIB and RF, not MN blocks sCD4-gp120, not as potent as MAb 15e - V2 binding MAbs BAT085 and G3-136 block G3-4 gp120 binding – sensitive to reduction of gp120 by DTT. Ho et al. [1992]
  - G3-4: Binding is sensitive to removal of glycans by endo H 50% neutralization of 4/9 primary isolates - has conformational features. Ho et al. [1991a]

No. 400

**MAb ID** BAT085 (BAT-085)

HXB2 Location gp160 (171–180)

Author Location gp120 (170-180 IIIB)

Epitope KEYAFFYKLD

Neutralizing L

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade IIIB HIV component: HIV-1

**Species (Isotype)** mouse (IgG1)

Research Contact Tanox Biosystems Inc and David Ho, ADARC, NY

References Parren et al. 1998a; Ditzel et al. 1997; Binley et al. 1997a; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; Wu et al. 1995; Yoshiyama et al. 1994; Gorny et al. 1994; Moore et al. 1994d; D'Souza et al. 1994; Moore et al. 1993a; Thali et al. 1993; Pirofski et al. 1993; Moore & Ho 1993; Fung et al. 1992; Fung et al. 1987

 BAT085: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated - authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]

- BAT085: Binding is blocked by other V2 region antibodies, enhanced by several anti-C1 MAbs, and anti-V3 MAb G511 – reciprocal enhancement of CD4i MAb 48d binding. Moore & Sodroski [1996]
- BAT085: Epitope suggested to be QKEYAFFYKLD binds oligomer – binding of V2 MAbs G3-136, G3-4 or BAT123 did not significantly alter gp120 dissociation from virus or expose the gp41 epitope of MAb 50-69, in contrast to anti-V3 MAbs. Poignard et al. [1996a]
- BAT085: Bound preferentially to the monomeric rather than oligomeric form of LAI gp120 – neutralizes cell free Hx10. Sattentau & Moore [1995]
- BAT085: HXB10 strain specificity binds native, deglycosylated, or denatured gp120. Wu et al. [1995]
- BAT085: Multi-lab study for antibody characterization and assay comparison – did not bind MN or SF2. D'Souza *et al*. [1994]
- BAT085: Interacts with two overlapping peptides with region of overlap KEYAFFYKLD. Gorny et al. [1994]
- BAT085: Neutralizes RF substitution 177 Y/H in the V2 loop of RF does not inhibit neutralization, in contrast to MAbs G3-4 and SC258. Yoshiyama *et al.* [1994]
- BAT085: Called BAT-85 conformational, does not bind well to denatured gp120 not reactive with SF-2 gp120, and does not inhibit HIV-1 sera from binding to IIIB gp120. Moore & Ho [1993]
- BAT085: 7/8 V2 murine MAbs required gp120 native structure to bind, but BAT085 was the exception type-specific. Moore *et al.* [1993a]
- BAT085: Peptide affinities of G3-136 and G3-4 are 100-fold less than BAT085, but BAT085 has lower affinity for BH10 gp120 and is weaker at neutralization. Moore *et al.* [1993a]
- BAT085: V2 region sCD4 does not block neutralizes IIIB and some primary isolates, but not MN or RF – binds MN – deglycosylation or DDT reduction of gp120 does not diminish reactivity. Fung *et al.* [1992]

**No.** 401

MAb ID 60b

**HXB2 Location** gp160 (172–181)

Author Location gp120 (172–181 HXB2)

Epitope EYAFFYKLDI

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2b)

References Shotton et al. 1995

 60b: V2 MAb did not neutralize HXB2 – bound to rgp120 in ELISA – substitutions 179-180 LD/DL and 191-193 YSL/GSS abrogate binding, as do changes outside the minimum epitope – competes with 12b, but not 74. Shotton *et al.* [1995]

**No.** 402

MAb ID 74

**HXB2 Location** gp160 (172–181)

**Author Location** gp120 (172–181)

Epitope EYAFFYKLDI

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG1)

References Shotton et al. 1995

• 74: V2 MAb did not neutralize HXB2 – did not bind rgp120 ELISA – position 179-180 LD to DL abrogates binding, as do changes outside the minimum epitope – does not compete with 60b or 12b, and is enhanced by two conformation dependent MAbs. Shotton *et al.* [1995]

No. 403

MAb ID 38/12b

HXB2 Location gp160 (172–191)

Author Location gp120 (172–191 HXB2)

Epitope EYAFFYKLDIIPIDNDTTSY

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat

References Wu et al. 1995

 38/12b: Broad specificity: HXB2, MN, SF162 – binds native and deglycosylated gp120. Wu et al. [1995]

**No.** 404

**MAb ID** 38/60b

**HXB2 Location** gp160 (172–191)

Author Location gp120 (172–191 HXB2)

Epitope EYAFFYKLDIIPIDNDTTSY

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat

References Wu et al. 1995

• 38/60b: Strain specificity: HXB2 – binds native and deglycosylated gp120. Wu *et al.* [1995]

**No.** 405

MAb ID polyclonal (VEI2)

**HXB2 Location** gp160 (176–196)

**Author Location** Env

Epitope FYKLDIVPIDNTTTSYRLISC

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Carlos et al. 1999

Antibody response to the epitopes in a vaccine construct (VEI) containing peptides from 5 hypervariable regions of gp120 was detected in the sera of HIV-1 positive subjects, including sera from 6 non-subtype B infections – serum samples from San Francisco, Canada and Puerto Rico cohort showed presence of antibodies against all five VEI hypervariable regions, but most consistently against the V3 region peptide NNNTRKSIRIGP-GRAFYTTGDIGNIRQ. Carlos et al. [1999]

**No.** 406

**MAb ID** 322-151 **Author Location** gp120 (200–217) **HXB2 Location** gp160 (211–221) Epitope PIPIHYCAPA Author Location gp120 (201–220 LAI) Neutralizing no Epitope EPIPIHYCAPA Immunogen vaccine Subtype B *Vector/Type:* protein *HIV component:* Env **Neutralizing** Species (Isotype) human References Valenzuela et al. 1998; Pincus et al. 1996; Immunogen vaccine Vector/Type: protein HIV component: Env Pincus & McClure 1993 Species (Isotype) mouse (IgG) • 110.1: There is another antibody with this ID that binds to Env Research Contact G. Robey, Abbot Labs at positions 491-500 in LAI, see. References Moore et al. 1994d; Moore et al. 1994c • 110.1: A panel of immunotoxins were generated by linking • 322-151: The relative affinity denatured/native gp120 is 30. Env MAbs to ricin A – immunotoxins mediated cell killing, but killing was not directly proportional to binding – 110.1-RAC Moore *et al.* [1994c] did not mediate cell killing, and sCD4 has no effect. Pincus & **No.** 407 McClure [1993]; Pincus et al. [1996] MAb ID 3D3.B8 HXB2 Location gp160 (211–221) No. 411 Author Location gp120 (211-220 LAI) MAb ID GV4H3 Epitope EPIPIHYCAPA HXB2 Location gp160 (219-226) Subtype B Author Location gp120 (219-226 IIIB) **Neutralizing** Epitope APAGFAIL Immunogen vaccine Neutralizing Vector/Type: protein HIV component: Env Immunogen vaccine Species (Isotype) mouse (IgG) Vector/Type: protein-Ab complex HIV com-References Moore et al. 1994c; Bolmstedt et al. 1990 ponent: gp120-Mab complex • 3D3.B8: The relative affinity denatured/native gp120 is greater Species (Isotype) mouse than 10. Moore et al. [1994c] References Denisova et al. 1996 • GV4H3: When anti-V3 MAb M77 was bound to gp120 and No. 408 used as an immunogen, it stimulated many MAbs to linear MAb ID 4C11.D8 epitopes. Denisova et al. [1996] **HXB2 Location** gp160 (211–221) Author Location gp120 (211–220 LAI) **No.** 412 Epitope EPIPIHYCAPA MAb ID J1 Subtype B HXB2 Location gp160 (222-231) Author Location gp120 (222-231 LAI) Neutralizing Epitope GFAILKCNNK Immunogen vaccine Vector/Type: protein HIV component: Env Subtype B Species (Isotype) mouse (IgM) Neutralizing References Moore et al. 1994c; Bolmstedt et al. 1990 Immunogen vaccine • 4C11.D8: The relative affinity denatured/native gp120 is Vector/Type: peptide Strain: B clade LAI greater than 10. Moore et al. [1994c] **Species (Isotype)** mouse (IgG1) Research Contact J. Hoxie, U. Penn. No. 409 References Cook et al. 1994; Moore et al. 1994d; Moore MAb ID 493-156 et al. 1994c HXB2 Location gp160 (211-230) • J1: MAbs against the glycosphingolipid GalCer block HIV Author Location gp120 (211-230 LAI) infection of normally susceptible CD4 negative cells from the Epitope EPIPIHYCAPAGFAILKCNN brain and colon – MAbs against the N-terminal half of gp120 Subtype B do not inhibit gp120 binding to GalCer - binding of GalCer to Neutralizing gp120 does not inhibit MAb binding. Cook et al. [1994] Immunogen vaccine • J1: The relative affinity denatured/native gp120 is 30. Moore Vector/Type: protein HIV component: Env et al. [1994c] Species (Isotype) mouse (IgG) Research Contact G. Robey, Abbot Labs No. 413 MAb ID J3 References Moore et al. 1994c • 493-156: The relative affinity denatured/native gp120 is >10. HXB2 Location gp160 (222–231) Moore et al. [1994c] Author Location gp120 (222-231 LAI) Epitope GFAILKCNNK No. 410 Subtype B **MAb ID** 110.1 Neutralizing HXB2 Location gp160 (212–221) Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI

Species (Isotype) mouse (IgG1)

Research Contact J. Hoxie, U. Penn.

References Cook et al. 1994; Moore et al. 1994c

• J3: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon – MAbs against the N-terminal half of gp120 do not inhibit gp120 binding to GalCer – binding of GalCer to gp120 does not inhibit MAb binding. Cook *et al.* [1994]

• J3: The relative affinity denatured/native gp120 is 30. Moore et al. [1994c]

**No.** 414

**MAb ID** 1006-30-D

HXB2 Location gp160 (236-245)

Author Location gp120 (241-251)

**Epitope** KGSCKNVSTV

Neutralizing

Immunogen

**Species** (**Isotype**) human ( $IgG1\lambda$ )

Ab Type gp120 C2

References Nyambi et al. 2000; Hioe et al. 2000

- 1006-30-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses CD4BS MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells C2 MAbs 1006-30-D and 847-D did not effect proliferation. Hioe *et al.* [2000]
- 847-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including two C2 MAbs the binding of anti-C2 MAbs was weak to isolates from clades B, C, D, E, F, G, and they did not not bind to isolates from subtypes A and H epitope is suggested to be in a 20 amino acid peptide KGSCKNVSTVQCTHGIRPVV. Nyambi *et al.* [2000]

**No.** 415

**MAb ID** 847-D

HXB2 Location gp160 (236–245)

**Author Location** gp120 (241–251)

**Epitope** KGSCKNVSTV

Neutralizing

Immunogen

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 C2

References Nyambi et al. 2000; Hioe et al. 2000

- 847-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses CD4BS MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells C2 MAbs 1006-30-D and 847-D did not effect proliferation. Hioe *et al.* [2000]
- 847-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including two C2 MAbs the binding of anti-C2 MAbs was weak to isolates from clades B, C, D, E, F, G, and they did not not bind to isolates from subtypes A and H epitope is suggested to be in a 20 amino acid peptide KGSCKNVSTVQCTHGIRPVV. Nyambi *et al.* [2000]

No. 416 MAb ID MF169.1 HXB2 Location gp160 (252–261)

Author Location gp120 (242–261 LAI)

Epitope RPVVSTQLLL

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

**Species (Isotype)** mouse (IgG)

References Moore et al. 1994d; Moore et al. 1994c; Thiri-

art et al. 1989

MF169.1: The relative affinity denatured/native gp120 is 11

 mutations 252 R/W, 257 T/G, and 257 T/R impair binding.
 Moore et al. [1994c]

No. 417

**MAb ID** MF170.1

HXB2 Location gp160 (252–261)

Author Location gp120 (242–261 LAI)

Epitope RPVVSTQLLL

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

References Moore et al. 1994d; Moore et al. 1994c; Thiriart et al. 1989

• MF170.1: The relative affinity denatured/native gp120 is 15 – mutations 252 R/W, 257 T/G, and 257 T/R impair binding to denatured and native gp120, and 262N/T, 269 E/L and 281 A/V to only native gp120. Moore *et al.* [1994c]

No. 418

MAb ID MF87.1

 $\textbf{HXB2 Location} \hspace{0.1cm} gp160 \hspace{0.1cm} (252\text{--}261)$ 

Author Location gp120 (242-261 LAI)

Epitope RPVVSTQLLL

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

References Moore et al. 1994c; Thiriart et al. 1989

MF87.1: The relative affinity denatured/native gp120 is 10

 mutations 252 R/W, 257 T/G, and 257 T/R impair binding.
 Moore *et al.* [1994c]

**No.** 419

**MAb ID** 213.1

**HXB2 Location** gp160 (252–261)

Author Location gp120 (242-261 LAI)

**Epitope** RPVVSTQLLL

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) mouse (IgG1)

Ab Type gp120 C2

Research Contact Claudine Bruck

**References** Moore *et al.* 1994c; Moore & Ho 1993; Thiriart *et al.* 1989

• 213.1: UK Medical Research Council AIDS reagent: ARP334.

• 213.1: The relative affinity denatured/native gp120 is 100 – mutations 252 R/W, 257 T/G or T/R impair binding. Moore et al. [1994c]

• 213.1: Bound preferentially to denatured IIIB and SF2 gp120. Moore & Ho [1993]

**No.** 420

MAb ID B12

HXB2 Location gp160 (252-271) Author Location gp120 (252-271 LAI)

Epitope RPVVSTQLLLNGSLAEEEVV

Subtype B

Neutralizing

Immunogen vaccine

HIV component: gp160

Species (Isotype) mouse (IgG)

Ab Type gp120 C2

References Maksiutov et al. 2002; Moore et al. 1994c

- B12: This epitope is similar to a fragment of human protein lymphatic endothelium-specific hyaluronan receptor LYVE-1, TTRLLVQGSLRAEE. Maksiutov et al. [2002]
- B12: C2 region the relative affinity for denatured/native gp120 is 27 - mutations 257 T/R and 262 N/T impair binding. Moore et al. [1994c]

**No.** 421

MAb ID B13 (Bh13, Chessie B13)

**HXB2 Location** gp160 (252–271)

Author Location gp120 (252-271 LAI)

Epitope RPVVSTQLLLNGSLAEEEVV

Subtype B **Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI HIV component: gp160

Species (Isotype) mouse (IgG2a)

Ab Type gp120 C2

Research Contact George Lewis, Institute of Human Virology,

Baltimore MD, USA

References Maksiutov et al. 2002; Wang et al. 2002c; Connor et al. 1998; Pincus et al. 1996; Moore et al. 1994d; Abacioglu et al. 1994; Moore et al. 1994c; Moore & Ho 1993; Pincus & McClure 1993

- B13: This epitope is similar to a fragment of human protein lymphatic endothelium-specific hyaluronan receptor LYVE-1, TTRLLVQGSLRAEE. Maksiutov et al. [2002]
- B13: Called Bh13 binds to gp120 but not to infected cells when linked to ricin A, the immunotoxin did not mediate cell killing - sCD4 has no effect. Pincus & McClure [1993]; Pincus et al. [1996]
- B13: C2 region epitope boundaries mapped by peptide scanning, core epitope: TQLLLN. Abacioglu et al. [1994]
- B13: The relative affinity for denatured/native gp120 is 30 mutations 257 T/R and 269 E/L impair binding. Moore et al.
- B13: Bound preferentially to denatured IIIB gp120. Moore & Ho [1993]

No. 422

MAb ID C13

**HXB2 Location** gp160 (252–271)

Author Location gp120 (252-271 LAI)

Epitope RPVVSTQLLLNGSLAEEEVV

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG1)

Ab Type gp120 C2

Research Contact George Lewis

References Maksiutov et al. 2002; Abacioglu et al. 1994; Moore et al. 1994c; Moore & Ho 1993

- Vector/Type: protein Strain: B clade LAI C13: NIH AIDS Research and Reference Reagent Program: 1209.
  - C13: This epitope is similar to a fragment of human protein lymphatic endothelium-specific hyaluronan receptor LYVE-1, TTRLLVQGSLRAEE. Maksiutov et al. [2002]
  - Epitope boundary extended to RPVVSTQLLL-NGSLAEEEVVIR, to take into account the effect of a point mutation. Abacioglu et al. [1994]
  - C13: The relative affinity for denatured/native gp120 is 36 - mutations 257 T/R, 267 E/L, and 269 E/L impair binding. Moore *et al.* [1994c]
  - C13: Bound preferentially to denatured IIIB gp120. Moore & Ho [1993]

No. 423

MAb ID M89

HXB2 Location gp160 (252-271)

Author Location gp120 (252-271 LAI)

Epitope RPVVSTQLLLNGSLAEEEVV

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) mouse (IgG1)

Ab Type gp120 C2

Research Contact Fulvia di Marzo Veronese

References Maksiutov et al. 2002; Moore et al. 1994d; Moore et al. 1994c; di Marzo Veronese et al.

- M89: This epitope is similar to a fragment of human protein lymphatic endothelium-specific hyaluronan receptor LYVE-1, TTRLLVQGSLRAEE. Maksiutov et al. [2002]
- M89: C2 region the relative affinity for denatured/native gp120 is >30 - mutations 257 T/R and 269 E/L impair binding. Moore *et al.* [1994c]
- M89: Immunoblot reactive, RIP negative, for strains IIIB, 451, MN, RF, and RUTZ. di Marzo Veronese et al. [1992]

No. 424

MAb ID B21

HXB2 Location gp160 (257–262)

**Author Location** gp120 (257–262 BH10)

Epitope TQLLLN

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI Vector/Type: protein Strain: B clade LAI HIV component: gp160 HIV component: gp160 Species (Isotype) mouse (IgG1) Species (Isotype) mouse (IgG1) Ab Type gp120 C2 Ab Type gp120 C2 References Abacioglu et al. 1994 References Abacioglu et al. 1994 • B21: C2 region, epitope boundaries mapped by peptide scan-• B3: C2 region, epitope boundaries mapped by peptide scanning. ning. Abacioglu et al. [1994] Abacioglu et al. [1994] No. 425 No. 429 MAb ID B23 MAb ID B26 HXB2 Location gp160 (257-263) HXB2 Location gp160 (257–262) **Author Location** gp120 (257–262 BH10) **Author Location** gp120 (257–263 BH10) **Epitope TQLLLN Epitope TQLLLNG** Neutralizing Neutralizing Immunogen vaccine Immunogen vaccine Vector/Type: protein Strain: B clade LAI Vector/Type: protein Strain: B clade LAI HIV component: gp160 HIV component: gp160 Species (Isotype) mouse (IgG2a) Species (Isotype) mouse (IgG1) Ab Type gp120 C2 Ab Type gp120 C2 References Abacioglu et al. 1994 References Abacioglu et al. 1994 • B23: C2 region, epitope boundaries mapped by peptide scanning. Abacioglu et al. [1994] ning. Abacioglu et al. [1994] **No.** 426 **No.** 430 MAb ID B24 MAb ID B29 HXB2 Location gp160 (257–262) HXB2 Location gp160 (257–263) **Author Location** gp120 (257–262 BH10) **Author Location** gp120 (257–263 BH10) **Epitope TQLLLN Epitope TQLLLNG Neutralizing Neutralizing** Immunogen vaccine Immunogen vaccine Vector/Type: protein Strain: B clade LAI Vector/Type: protein Strain: B clade LAI HIV component: gp160 HIV component: gp160 Species (Isotype) mouse (IgG2a) Species (Isotype) mouse (IgG2a) Ab Type gp120 C2 Ab Type gp120 C2 References Abacioglu et al. 1994 References Abacioglu et al. 1994 • B24: C2 region, epitope boundaries mapped by peptide scanning. Abacioglu et al. [1994] ning. Abacioglu et al. [1994] No. 427 **No.** 431 MAb ID B25 MAb ID B36 HXB2 Location gp160 (257–262) HXB2 Location gp160 (257–263) **Author Location** gp120 (257–262 BH10) **Author Location** gp120 (257–263 BH10) Epitope TQLLLN Epitope TQLLLNG Neutralizing **Neutralizing** Immunogen vaccine Immunogen vaccine Vector/Type: protein Strain: B clade LAI Vector/Type: protein Strain: B clade LAI HIV component: gp160 HIV component: gp160 **Species (Isotype)** mouse (IgG1) Species (Isotype) mouse (IgG1) Ab Type gp120 C2 Ab Type gp120 C2 References Abacioglu et al. 1994 References Abacioglu et al. 1994 • B25: C2 region, epitope boundaries mapped by peptide scanning. Abacioglu et al. [1994] ning. Abacioglu et al. [1994] No. 428 No. 432 MAb ID B3 **MAb ID** 110.E **HXB2 Location** gp160 (257–262) **HXB2 Location** gp160 (262–281) **Author Location** gp120 (257–262 BH10) Author Location gp120 (262-281 LAI) Epitope NGSLAEEEVVIRSVNFTDNA Epitope TQLLLN

Subtype B

**Neutralizing** 

gp160 Antibodies

Neutralizing Immunogen vaccine

**HIV Antibodies Tables** 

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C2

Research Contact F. Traincard

References Maksiutov et al. 2002; Moore et al. 1994d; Moore et al. 1994c

- 110.E: This epitope is similar to a fragment of human protein lymphatic endothelium-specific hyaluronan receptor LYVE-1, TTRLLVQGSLRAEE. Maksiutov et al. [2002]
- 110.E: The relative affinity for denatured/native gp120 is 7.3. IIIB-V3-21: The role of serine proteases on HIV infection was Moore *et al.* [1994c]

**No.** 433

**MAb ID** 110.C

HXB2 Location gp160 (271-280) Author Location gp120 (271–280 LAI)

**Epitope** VIRSVNFTDN

Subtype B

**Neutralizing** 

Immunogen vaccine

HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C2

Research Contact F. Traincard, Hybridolabs, Institut Pasteur References Valenzuela et al. 1998; Moore et al. 1994d; Moore et al. 1994c

- 110.C: Only slightly reduces LAI viral binding or entry into CEM cells. Valenzuela et al. [1998]
- 110.C: The relative affinity for denatured/native gp120 is 1. Moore et al. [1994c]

No. 434

MAb ID IIIB-V3-26

HXB2 Location gp160 (291–307)

Author Location gp120 (299–304 IIIB)

Epitope SVEINCTRPNNNTRKSI

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

References Maksiutov et al. 2002; Laman et al. 1992

- IIIB-V3-26: This epitope is similar to a fragment of the FasI receptor precursor (Apptosis-mediating surface antigen fas) (APO-1 antigen) (CD95 antigen), VEINCTRQN. Maksiutov et al. [2002]
- IIIB-V3-26: Binds to the base of the V3 loop on denatured gp120. Laman et al. [1992]

No. 435

**MAb ID** IIIB-V3-21 (V3-21)

HXB2 Location gp160 (294-299)

Author Location gp120 (299-304 IIIB)

**Epitope INCTRP** 

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

**Species (Isotype)** mouse (IgG1)

Ab Type gp120 V3

Research Contact J. Laman

References Ling et al. 2004; Maksiutov et al. 2002; Zhang et al. 2002; Valenzuela et al. 1998; Laman et al. 1993; Laman et al. 1992

**Keywords** antibody binding site definition and exposure

- IIIB-V3-21: UK Medical Research Council AIDS reagent: ARP3048.
- IIIB-V3-21: NIH AIDS Research and Reference Reagent Program: 1725.
- explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of the V3 MAbs 694-98D and 447-52D, that both bind near the tip of the loop, was decreased by both thrombin and trypsin, but anti-V3 MAb IIIB-V3-21 was not decreased in either case. Ling et al. [2004] (antibody binding site definition and exposure)
- Vector/Type: protein Strain: B clade LAI IIIB-V3-21: This epitope is similar to a fragment of the FasI receptor precursor (Apptosis-mediating surface antigen fas) (APO-1 antigen) (CD95 antigen), VEINCTRQN. Maksiutov
  - IIIB-V3-21: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera - 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 – thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002]
  - IIIB-V3-21: Does not block HIV-1 LAI binding or entry into CEM cells. Valenzuela et al. [1998]
  - IIIB-V3-21: Binds to NP40 treated gp120, and epitope is probably obscured by local glycosylation. Laman et al. [1993]
  - IIIB-V3-21: Binds to the base of the V3 loop on denatured gp120. Laman et al. [1992]

No. 436

**MAb ID** 168B8

HXB2 Location gp160 (296-317)

Author Location gp120 (BaL)

Epitope CTRPNYNKRKHIGPGRAF

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: gp120-CD4 complex HIV component: gp120 Adjuvant: Ribi adjuvant

(MPL+TDM) (RIBI)

**Species (Isotype)** humanized mouse (IgG2 $\kappa$ )

**Ab Type** gp120 V3

Research Contact Abraham Pinter, Lab. of Retrovirology, Public Research Institute, pinter@phri.org

References He et al. 2003

Keywords antibody binding site definition and exposure, Species (Isotype) human (IgM) vaccine antigen design

• 168B8: Vaccination of a gp120-CD4 fusion complex in six transgenic XMG2 XenoMouse mice that produce human IgG2 with K light chain did not produce any neutralizing antibodies. 36/39 MAbs derived from one of these mice were in one of two competition groups that were conformational and specific for the complex, suggesting this chimeric vaccine may be of little value, as immunodominant responses are to epitopes unique to the complex, not present in native Env. This MAb is one of the 3/36 non-neutralizing MAbs that bound to linear epitopes on gp120. He et al. [2003] (antibody binding site definition and exposure, vaccine antigen design)

No. 437

MAb ID polyclonal

HXB2 Location gp160 (296-331)

Author Location gp120 (MN)

Epitope CNYNKRKRIHIGPGRAFYTTKNIIGTIC

**Neutralizing** L Immunogen

Species (Isotype) rabbit (IgA, IgG)

Ab Type gp120 V3

References FitzGerald et al. 1998

• Polyclonal response to MN, or Thai E V3 loop inserted into Pseudomonas Exotoxin for vaccination - inserts of 14 or 26 amino acids were used from MN or a Thai E strain, constrained by disulfide bond – sera from vaccinated rabbit were reactive with strain-specific gp120 – administration to mucosal surfaces elicits IgA. FitzGerald et al. [1998]

No. 438

MAb ID polyclonal

HXB2 Location gp160 (297–330)

Author Location Env (303-335 LAI)

Epitope TRPNNNTRKSIHIGPGRAFYATGEIIGDIRQAH

Subtype B Neutralizing no Immunogen vaccine

> Vector/Type: lipopeptide Strain: B clade LAI HIV component: V3 Adjuvant: QS21

Species (Isotype) human (IgG)

**Ab Type** gp120 V3

References Pialoux et al. 2001

• 28 subjects were vaccinated with six HIV-1 peptides that were selected to be particularly rich in CTL epitopes, presented in lipopeptides with or without adjuvant QS21 - HIV-specific Ab responses were detected in 25/28 (89%), proliferative in 19/28 (79%), and CTL in 13/24 (54%) of testable volunteers – 14/28 had non-neutralizing Ab responses to this peptide (E), 7/24 had proliferative responses, and multiple CTL responses were detected. Pialoux et al. [2001]

**No.** 439

MAb ID MO97/V3

HXB2 Location gp160 (299-308)

Author Location gp120 (299–308 IIIB)

Epitope PNNNTRKSIR

Neutralizing no

Immunogen in vitro stimulation or selection

Ab Type gp120 V3

References Gorny & Zolla-Pazner 2004; Ohlin et al. 1992 Keywords review

- M097/V3: Review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains; a subset can also neutralize some primary isolates. The three IgMs, M096, M097, and M099, are nonneutralizing. Gorny & Zolla-Pazner [2004] (review)
- MO97: Generated through in vitro stimulation of uninfecteddonor lymphocytes with rpB1 (IIIB Env 286-467) Ohlin et al. [1992]

**No.** 440

MAb ID polyclonal

HXB2 Location gp160 (299-331)

**Author Location** gp120 (306–338 BH10)

Epitope PNNNTRKSIRIQRGPGRAFVTIGKIGNMRQAHC

Neutralizing L Immunogen vaccine

Vector/Type: peptide Strain: B clade BH10

Species (Isotype) rabbit (IgG)

Ab Type gp120 V3

References Neurath & Strick 1990

• 21 V3 loop variant peptides spanning this region were tested and serological cross-reactivity correlated with divergence. Neurath & Strick [1990]

No. 441

MAb ID 55/11

**HXB2 Location** gp160 (300–315)

**Author Location** gp120 (300–315)

Epitope NNNTRKRIRIQRGPGR?

**Neutralizing** 

Immunogen

Species (Isotype)

Ab Type gp120 V3

References Peet et al. 1998

• 55/11: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic – these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind, and anti-V3 MAb 55/11 binding was abrogated by V3 serine substitutions in the V3 loop - mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]

No. 442

**MAb ID** 8/38c (8/38/1c)

HXB2 Location gp160 (300-315)

**Author Location** gp120 (300–315 HXB10)

Epitope NNNTRKRIRIQRGPGR

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2a)

Ab Type gp120 V3

Research Contact C. Dean and C. Shotton, Institute for Cancer Author Location gp120 Research, Surrey, UK

References Peet et al. 1998; Parren et al. 1998a; Jeffs et al. 1996; Sattentau & Moore 1995; McKeating et al. 1992a

- 8/38c: UK Medical Research Council AIDS reagent: ARP3039.
- 8/38c: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated – authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- 8/38c: Called 8/38/1c: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic – these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind, and anti-V3 MAb 8/38c binding was only diminished by V3 serine substitutions C-term to the tip of the loop - mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]
- 8/38c: Deletion of the V1V2 regions did not affect anti-V3 Abs ability to bind when compared to intact rec gp120. Jeffs et al.
- 8/38c: Binds equally well to monomer and oligomer, less rapid association rate than other anti-V3 antibodies, and an associated less potent neutralization of lab strains. Sattentau & Moore [1995]
- 8/38c: Binds to virion gp120 and neutralizes only in the presence of sCD4. McKeating et al. [1992a]

**No.** 443

MAb ID 8/64b

HXB2 Location gp160 (300-315)

**Author Location** gp120 (300–315 HXB10)

Epitope NNNTRKRIRIQRGPGR

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10 HIV component: gp120

Species (Isotype) rat (IgM)

Ab Type gp120 V3

References Peet et al. 1998; McKeating et al. 1992a

- 8/64b: UK Medical Research Council AIDS reagent: ARP3036.
- 8/64b: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic - these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind, and anti-V3 MAb 8/64b binding was abrogated by V3 serine substitutions C-term to the tip of the loop – mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]
- 8/64b: Binds to virion gp120 and neutralizes only in the presence of sCD4. McKeating et al. [1992a]

No. 444 MAb ID polyclonal HXB2 Location gp160 (300-321)

Epitope NYNKRKRIHIGPGRAFYTTK

**Neutralizing** L

Immunogen HIV-1 infection, vaccine

Vector/Type: peptide HIV component: V3

Species (Isotype) human

Ab Type gp120 V3

References Bartlett et al. 1998

V3 peptide vaccine (MN, RF, EV91, and Can0A) with a C4 helper T cell epitope were used to vaccinate HLA-B7 HIVinfected patients – V3 Ab levels and the anti-HIV proliferative response, but no decrease in HIV-1 RNA levels or increase in CD4 levels was observed. Bartlett et al. [1998]

No. 445

MAb ID polyclonal

HXB2 Location gp160 (300–321)

**Author Location** gp120

Epitope NYNKRKRIHIGPGRAFYTTK

**Neutralizing** 

Immunogen HIV-1 exposed seronegative

Species (Isotype) human (IgA)

Ab Type gp120 V3

References Kaul et al. 1999

· HIV-1 Env-specific mucosal IgA found in genital track of 16/21 HIV-1 resistant chronically exposed Kenyan sex workers - 11/21 had detectable Th responses. Kaul et al. [1999]

No. 446

MAb ID polyclonal

HXB2 Location gp160 (300–322)

**Author Location** gp120 (IIIB)

Epitope CNNTRKSIRIQRGPGRAFVTIGK

**Neutralizing** L

**Immunogen** 

**Species** (**Isotype**) guinea pig (IgG)

**Ab Type** gp120 V3

Research Contact D. Bolognesi and T. Matthews, Duke Univer-

References Allaway et al. 1993

• Synergy with combinations of CD4-based molecules in inhibition of HIV-1 Env mediated cell fusion. Allaway et al. [1993]

No. 447

MAb ID polyclonal (VEI3)

HXB2 Location gp160 (300-328)

**Author Location** Env

Epitope NNNTRKSIRIGPGRAFYTTGDIGNIRQ

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Carlos et al. 1999

• Antibody response to the epitopes in a vaccine construct (VEI) containing peptides from 5 hypervariable regions of gp120 was detected in the sera of HIV-1 positive subjects, including sera from 6 non-subtype B infections - serum samples from San

Francisco, Canada and Puerto Rico cohort showed presence of antibodies against all five VEI hypervariable regions, but most

GRAFYTTGDIGNIRQ. Carlos et al. [1999]

No. 448

**MAb ID** 9284 (NEA 9284)

HXB2 Location gp160 (301-312)

Author Location gp120 (307–318 IIIB)

Epitope NNTRKSIRIQRG

Neutralizing L Immunogen vaccine

Vector/Type: inactivated HIV Strain: B

clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

Research Contact Dupont de Nemours, Les Ulis, France or

Wilmington, Delaware

References Schonning et al. 1998; Parren et al. 1998a; Binley et al. 1997a; Cao et al. 1997b; Poignard et al. 1996a; Moore & Sodroski 1996; Fontenot et al. 1995; VanCott et al. 1995: Sattentau & Moore 1995: Sorensen et al. 1994; Okada et al. 1994; Cook et al. 1994; Thali et al. 1994; VanCott et al. 1994; Thali et al. 1993; Trujillo et al. 1993; Moore et al. 1993b; Sattentau et al. 1993; McKeating et al. 1992a; Wyatt et al. 1992; Sattentau & Moore 1991; Skinner et al. 1988a; Skinner et al. 1988b

- 9284: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated - authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- 9284: In a study of the influence of the glycan at position 306 of the V3 loop on MAb recognition, 9284 was found to have an inaccessible epitope on the oligomeric form of Env and anti-V3 MAbs were found to neutralize an HIV-BRU mutant virus that lacks the V3 loop glycan more efficiently than HIV-BRU. Schonning et al. [1998]
- 9284: Virus with the V1-V2 loop deleted was viable and more susceptible to neutralization by CD4i MAb 17b, and anti-V3 MAbs 1121, 9284, and 110.4, but not to and CD4BS MAb F105 or sCD4. Cao et al. [1997b]
- 9284: Binds V3 loop anti-C1 MAbs 133/290 and 135/9 enhance binding - reciprocal binding inhibition of other anti-V3 MAbs. Moore & Sodroski [1996]
- 9284: V3 MAbs 9284, BAT123, 110.5, and 110.I could each significantly increase gp120 dissociation from virus, mimicking sCD4, and expose the gp41 epitope for MAb 50-69, in contrast to anti-V2 MAbs. Poignard et al. [1996a]
- 9284: Binds equally well to monomer and oligomer, rapid association and potent neutralization of lab strains – neutralizes cell-free virus Hx10. Sattentau & Moore [1995]
- 9284: Used to monitor HIV-1 Env expression in infected H9 cells, binds native and reduced gp120s similarly. VanCott et al. [1995]
- 9284: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon – this MAb can inhibit gp120 binding to GalCer in vitro. Cook et al. [1994]

consistently against the V3 region peptide NNNTRKSIRIGP- • 9284: Binding domain aa 301-310: TRKSIRIORG – mutations in the V3 loop from basic residues can destroy virus infectivity and syncytium formation: R306T, R309T and R313G can also reduce binding of V3 MAbs with two different binding sites: 9284 and 0.5beta - called NEA9284. Okada et al. [1994]

- 9284: Did not neutralize infection of HIV/HTLV-I pseudotype. Sorensen et al. [1994]
- 9284: gp41 mutation that confers resistance to neutralization by anti-CD4 binding site antibodies does not reduce neutralizing efficiency of this V3 region MAb. Thali et al. [1994]
- 9284: Does not bind MN gp120, just IIIB. VanCott et al. [1994]
- 9284: Inhibits C4 region antibodies (G3-299, G3-519) which have conformational requirements. Moore et al. [1993b]
- 9284: Increased binding in the presence of sCD4. Sattentau et al. [1993]
- 9284: Peptide RIQRGPGRAFVTIGKIGNMRQA Reacts with three human brain proteins of 35, 55, 110 kd - called NEA-9284. Trujillo et al. [1993]
- 9284: Single amino acid substitutions in the C4 region (427 W/V or W/S) or at the base of the V3 loop (298 R/G) can significantly increase binding and neutralization-position 427 is also important for CD4 binding and anti-CD4 binding site MAbs. Wyatt et al. [1992]
- 9284: Two fold increase in binding to gp120 in the presence of bound sCD4. Sattentau & Moore [1991]
- 9284: IIIB type-specific binding and neutralization. Skinner et al. [1988b]

No. 449

MAb ID polyclonal

HXB2 Location gp160 (301-325)

Author Location gp120 (IIIB)

Epitope NNTRKSIRIQRGPGRAFVTIGKIGN

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

Adjuvant: Cholera toxin (CT)

Species (Isotype) mouse (IgA)

Ab Type gp120 V3

References Bukawa et al. 1995

Polyclonal secretory IgA antibody raised by mucosal immunization is able to neutralize IIIB, SF2, and MN - HIV-1 neutralization may be due to V3, CD4 or HPG30 component of the multicomponent peptide immunogen. Bukawa et al. [1995]

No. 450

MAb ID polyclonal

HXB2 Location gp160 (301–325)

Author Location gp120 (IIIB)

Epitope NNTRKSIRIQRGPGRAFVTIGKIGN

Neutralizing L

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB

HIV component: Env, Rev

**Species (Isotype)** mouse (IgA22a)

Ab Type gp120 V3

References Sasaki et al. 1998

 An anti-env response was sought, and co-expression of Rev was required – intramuscular versus nasal vaccination with DNA vaccine with a QS21 adjuvant was studied – QS21 enhanced the IgG2a response mediated via Th1 cytokines IFNgamma and IL-2. Sasaki et al. [1998]

No. 451

MAb ID polyclonal

HXB2 Location gp160 (302-317)

Author Location Env (B consensus)

Epitope NTRKSIHIGPGRAF

Subtype B

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Morris et al. 2001

Ab responses before HAART therapy and after one year of therapy were measured in 8 individuals that were classified HAART successes, and 10 patients who were classified as HAART failures – V3 peptide antibody binding titers to the B-consensus and MN and SF2 variants, and neutralization of HIV-1 MN and four subtype B clinical isolates were tested – subjects with strong anti-V3 and NAb humoral immune responses before starting HAART were more likely to achieve sustained viral suppression to <500 copies RNA/ml on HAART – HIV-specific Ab responses declined after 1 year of successful viral suppression on HAART. Morris et al. [2001]</li>

No. 452

MAb ID polyclonal

HXB2 Location gp160 (302–318)

**Author Location** Env

Epitope NTRKSIHIGPGRAFY

Neutralizing LP

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Bongertz et al. 2001

• Non-transmitting mothers had an increased frequency of high neutralizing plasma Ab titers against HIV-1 MN (1:50 dilution, >90% neutralization, 33/88 pregnant women), compared to plasma from transmitting mothers (0/8 pregnant women) – non-transmitting mothers also had more potent neutralization against primary isolates from transmitting mothers, but neutralization of autologous virus was comparable for non-transmitting (7/13) and transmitting mothers (2/4) Bongertz et al. [2001]

No. 453

MAb ID MAG 109

HXB2 Location gp160 (302-321)

**Author Location** gp120 (302–321 BH10)

Epitope NTRKSIRIQRGPGRAFVTIG

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain:

B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 V3

References Kang et al. 1994

• MAG 109: Binds a V3 loop peptide – sensitive to both V3 loop mutations and a mutation at the base of the V1/V2 loop structure (120/121 VK/LE) Kang *et al.* [1994]

**No.** 454

**MAb ID** MAG 49 (#49)

**HXB2 Location** gp160 (302–321)

**Author Location** gp120 (302–321 BH10)

Epitope NTRKSIRIQRGPGRAFVTIG

**Neutralizing** L

Immunogen vaccine

*Vector/Type:* sCD4-gp120 complex *Strain:* 

B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 V3

References Moore & Sodroski 1996; Kang et al. 1994

- MAG 49: Called #49 in this text. Binding enhanced by anti-C1 MAbs 133/290, 135/9, and by many anti-CD4 binding site MAbs – reciprocal enhancement of some anti-V2 MAbs – reciprocal binding inhibition of anti-V3 MAbs. Moore & Sodroski [1996]
- MAG 49: Binds a V3 loop peptide sensitive to both V3 loop mutations and a mutation at the base of the V1/V2 loop structure (120/121 VK/LE) Kang *et al.* [1994]

No. 455

MAb ID MAG 53

HXB2 Location gp160 (302–321)

**Author Location** gp120 (302–321 BH10)

Epitope NTRKSIRIQRGPGRAFVTIG

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain:

B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 V3

References Kang et al. 1994

 MAG 53: Binds a V3 loop peptide – sensitive to both V3 loop mutations and a mutation at the base of the V1/V2 loop structure (120/121 VK/LE) Kang et al. [1994]

**No.** 456

MAb ID MAG 56

HXB2 Location gp160 (302-321)

Author Location gp120 (302-321)

Epitope NTRKSIRIQRGPGRAFVTIG

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 V3

References Kang et al. 1994

 MAG 56: Binds a V3 loop peptide – sensitive to both V3 loop mutations and a mutation at the base of the V1/V2 loop structure (120/121 VK/LE) Kang et al. [1994]

No. 457

**MAb ID** 1324-E (1324E)

(Zol-

**HXB2 Location** gp160 (303–308) **Author Location** Env (subtype CRF01)

**Epitope** TRTSVR **Subtype** CRF01\_AE

Neutralizing L

Immunogen HIV-1 infection Species (Isotype) human (IgG1κ) Ab Type gp120 V3

Research Contact Susan Zolla-Pazner

las01@mcrcr6.med.nyu) (NYU Med.

Center)

**References** Gorny & Zolla-Pazner 2004; Nyambi *et al.* 2000; Zolla-Pazner *et al.* 1999b; Zolla-Pazner

et al. 1999a; Gorny et al. 1998

**Keywords** antibody generation, review, subtype comparisons, variant cross-recognition or cross-

neutralization

- 1324-E: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 1324-E: Called 1324E A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding V3 MAbs tended to have the most cross-reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H 1324E showed poor cross-reactivity, and was the only MAb tested that was derived from a non-B clade infected patient, an E clade infection was the source of 1324E. Nyambi *et al.* [2000] (**subtype comparisons**)
- 1324-E: E clade stimulated MAb did not cross-react with B clade peptides nor did B clade derived peptides with an E clade V3 loop, but both E and B clade stimulated Abs can cross-react with some peptides from other clades this Ab showed strong binding to several E, A and F peptides, one C peptide, and no reactivity with B peptides and most D peptides. Zolla-Pazner et al. [1999a] (subtype comparisons)
- 1324-E: MAb reacted with peptides from E clade, while B clade derived MAbs could not. Zolla-Pazner *et al.* [1999b] (**subtype comparisons**)
- 1324-E: A human MAb was derived from an HIV-1 E clade infection from a US service man who had served in Thailand, selected with the consensus V3 peptide from clade E cross-reactive with V3 peptides, and gp120 from E, C and A clades, as well as cells infected with a C-clade primary isolate, but not with B and D clade V3 peptides or rgp120 neutralizes E clade virus adapted for growth in H9 cells, but not 5 primary E clade isolates, including the autologous isolate kinetic parameters were measured, 1324E was comparable to 447-52D. Gorny et al. [1998] (antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

No. 458
MAb ID polyclonal
HXB2 Location gp160 (303–319)
Author Location gp120 (subtype C)
Epitope CKRKIHIGPGQAFYT
Subtype C
Neutralizing

Immunogen vaccine

Vector/Type: peptide in ISCOM, peptide in liposome HIV component: V3 Adjuvant: Immune stimulating complexes (ISCOM)

Species (Isotype) mouse (IgG2a, IgG2b)

Ab Type gp120 V3

References Ahluwalia et al. 1997

 A V3 loop peptide modified to resemble an Indian form (GPGQ) was incorporated into ISCOMS (immune stimulating complexes) or liposomes, and used to immunize mice – the IgG2a/IgG2b antibody response was enhanced by the presentation in the ISCOM suggestive of a Th1 response. Ahluwalia et al. [1997]

No. 459

MAb ID MO99/V3

HXB2 Location gp160 (304–308)

Author Location gp120 (304–308 IIIB)

Epitope RKSIR

Neutralizing no

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgM)

**Ab Type** gp120 V3

**References** Gorny & Zolla-Pazner 2004; Ohlin *et al.* 1992 **Keywords** antibody binding site definition and exposure, antibody generation

- M099/V3: Review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains; a subset can also neutralize some primary isolates. The three IgMs, M096, M097, and M099, are nonneutralizing. Gorny & Zolla-Pazner [2004]
- MO99: Generated through in vitro stimulation of uninfecteddonor lymphocytes with rpB1 (IIIB Env 286-467) Ohlin et al. [1992] (antibody binding site definition and exposure, antibody generation)

No. 460

MAb ID C311E

HXB2 Location gp160 (304–313)

Author Location gp120 (309–316 MN)

Epitope RKRIHIGP

**Neutralizing** L

Immunogen HIV-1 infection

**Species (Isotype)** chimpanzee (IgG1)

Ab Type gp120 V3

References Alsmadi & Tilley 1998; Warrier et al. 1996

- C311E: A study of 6 anti-Env MAbs and their ability to bind or direct ADCC against target cells infected with IIIB, MN, SF-2, and RF – C311E bound and directed lysis against all four strains. Alsmadi & Tilley [1998]
- C311E: Chimps were infected with HIV-1 IIIB, and this resulting MAb gave synergistic neutralization of HIV-1 when combined with anti-V2 MAb C108G. Warrier et al. [1996]

No. 461

**MAb ID** 907

**HXB2 Location** gp160 (304–314)

**Author Location** gp120 (309–318)

Epitope RKSIRIQRGPG

**Neutralizing** L

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: gp160

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

**References** Pincus *et al.* 1996; Pincus *et al.* 1991; Pincus *et al.* 1989; Chesebro & Wehrly 1988

- 907: A panel of immunotoxins were generated by linking Env MAbs to ricin A – immunotoxins mediated cell killing, but killing was not directly proportional to binding. Pincus *et al.* [1996]
- 907: Epitope sequence is based on database count of a specified location 924-RAC immunotoxin is IIIB strain-specific. Pincus *et al.* [1991]
- 907: Coupled to ricin A chain (RAC), MAb 907 inhibited protein synthesis and cell growth in HIV-infected cells. Pincus et al. [1989]
- 907: Strain specific binding, and neutralization of only the LAV strain. Chesebro & Wehrly [1988]

No. 462

**MAb ID** 924

**HXB2 Location** gp160 (304–314) **Author Location** gp120 (309–318 IIIB)

Epitope RKSIRIQRGPG

**Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type gp120 V3

**References** Pincus *et al.* 1998; Pincus *et al.* 1996; Cook *et al.* 1994; Pincus *et al.* 1993; Pincus & McClure 1993; Pincus *et al.* 1991; Chesebro & Wehrly 1988

- 924: A panel of immunotoxins were generated by linking Env MAbs to ricin A – immunotoxins mediated cell killing, but killing was not directly proportional to binding. Pincus *et al.* [1996]
- 924: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon – this MAb can inhibit gp120 binding to GalCer in vitro. Cook et al. [1994]
- 924: MAb was coupled to ricin A chain (RAC) immunotoxin efficacy was not significantly decreased by sCD4, although the efficacy of gp41 MAb immunotoxins *in vitro* increased 30-fold by sCD4. Pincus & McClure [1993]
- 924: Ab response in IIIB lab workers was compared to gp160 LAI vaccine recipients MAb 924 was used as a control infected lab workers and a vaccinia gp160 vaccine had strong V3 MAb response, but alum absorbed rec gp160 did not generate anti-V3 response. Pincus et al. [1993]
- 924: Epitope sequence is based on database count of a specified location 924-RAC immunotoxin is IIIB strain-specific. Pincus et al. [1991]
- 924: HIV IIIB strain specific. Chesebro & Wehrly [1988]

**No.** 463

MAb ID polyclonal

**HXB2 Location** gp160 (304–318)

**Author Location** gp120 (304–318 LAI) **Epitope** RKSIRIQRGPGRAFV Subtype B

Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgG, IgM)

Ab Type gp120 V3

References Chin et al. 1995

 Mimicking the humoral immune response in vitro supports isotype switching – human IgG MAbs were generated from naive donors. Chin et al. [1995]

No. 464

MAb ID polyclonal

**HXB2 Location** gp160 (304–318)

Author Location gp120 (304-318 LAI)

Epitope RKSIRIQRGPGRAFV

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI

Species (Isotype) human (IgG, IgM)

Ab Type gp120 V3

References Zafiropoulos et al. 1997

IgG to IgM isotype switching in response to primary and secondary peptide vaccinations was studied – the immunogen contained a V3 loop fragment and a tetanus toxin helper epitope. Zafiropoulos *et al.* [1997]

No. 465

MAb ID polyclonal

**HXB2 Location** gp160 (304–318)

Author Location gp120 (NY5)

Epitope KKGIAIGPGRTLY

Neutralizing

**Immunogen** 

Species (Isotype) (IgM)

Ab Type gp120 V3

References Metlas et al. 1999a; Metlas et al. 1999b

• Auto-Abs that react with the V3 loop of NY5 are present in the sera of HIV- individuals, and are predominantly IgM. Metlas *et al.* [1999b]

**No.** 466

**MAb ID** D19

HXB2 Location gp160 (304–320)

Author Location gp120 (V3) (MN)

**Epitope** RKRIHIGPGRAFYTT

Subtype A, B, F

Neutralizing yes

Immunogen vaccine

Vector/Type: protein Strain: B clade BH8

HIV component: gp140

Species (Isotype) mouse (IgG)

**Ab Type** gp120 CD4i, gp120 V3

Research Contact Paolo Lusso, Human Virology, San Raffaele Scientific Institute, Milan, Italy. paolo@hsr.it

References Lusso et al. 2005

Keywords antibody binding site definition and expo-

sure, antibody generation, subtype comparisons, variant cross-recognition or cross-

neutralization

• D19: The epitope for D19 is conserved and embedded in V3. Species (Isotype) human (IgG1 $\kappa$ ) D19 is unique because for R5 viruses, it was cryptic and did not bind without exposure to sCD4, but for X4 and R5X4 isolates it Research Contact Susan was constitutively exposed. It had a similar overlapping binding region with MAbs 447-52D, B4e8, and 268-D, but different reactivity patterns and fine specificity; D19 binding to monomeric gp120 was independent of sCD4, the dependence was only seen in the context of native oligomeric Env. D19 reacted with 23/29 B clade Envs, but to only 2/14 viruses from other clades: one A and one F, but no C, D or E clade strains. D19 can neutralize X4 and R5X4 isolates, but could only neutralize R5 isolates in the presence of sCD4. The authors suggest that a more exposed V3 loop may facilitate CXCR4 coreceptor usage, but that this phenotype is limited in vivo by neutralizing antibodies until the onset of progressive disease. Lusso et al. [2005] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

No. 467

**MAb ID** 10F10

HXB2 Location gp160 (304-320)

Author Location gp120 (MN)

Epitope RKRIHIGPGRAFYTT

Neutralizing L

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN

HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

References Duarte et al. 1994

• 2C4: Putative epitope lies within IHIGPGRAFYT – generated by multi-epitope polypeptide immunization - recognize MN and SC (TRSIHIGPGRAFYTT) peptides, lower affinity for SF2. Duarte et al. [1994]

No. 468

MAb ID 2C4

HXB2 Location gp160 (304-320)

Author Location gp120 (MN)

Epitope RKRIHIGPGRAFYTT

**Neutralizing** L (MN)

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN

Species (Isotype) mouse (IgG2a)

Ab Type gp120 V3

References Duarte et al. 1994

• 2C4: Putative epitope lies within IHIGPGRAFYT – neutralizes MN, not IIIB and SF2 – generated by multi-epitope polypeptide immunization – recognize MN and SC (TRSIHIGPGRAFYTT) peptides, lower affinity for SF2. Duarte et al. [1994]

No. 469

MAb ID 412-D (412-10D, 412, 412D)

HXB2 Location gp160 (304-320)

Author Location gp120 (MN)

Epitope RKRIHIGPGRAFYTT

Subtype B

**Neutralizing** L

Immunogen HIV-1 infection

Ab Type gp120 V3

Zolla-Pazner (Zol-(NYU las01@mcrcr6.med.nyu) Med.

Center)

References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Nyambi et al. 1998; Gorny et al. 1998; Fontenot et al. 1995; VanCott et al. 1994; Spear et al. 1993;

Gorny et al. 1993

Keywords antibody binding site definition and exposure, binding affinity, complement, kinetics, review, subtype comparisons, vaccine antigen design, variant cross-recognition or crossneutralization

- 412-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 412-D: Called 412: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 412-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H - 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding - V3 MAbs tended to have the most crossreactive binding to clade A, B, C, and D isolates, less to E, F, G, and H – 412-D showed limited reactivity. Nyambi et al. [2000] (subtype comparisons)
- 412-D: Review of clade specificity and anti-V3 HIV-1-Abs. Zolla-Pazner et al. [1999a] (review, subtype comparisons)
- 412-D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 391.5, 412 and 418, all selected with MN V3 peptide - the core amino acids HIGPGR tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure)
- 412-D: Kinetic parameters were measured, and the association rates were similar, but dissociation rate constants were quite variable for V3 MAbs, 412-D has a relatively fast dissociation, thus low affinity among V3 MAbs. Gorny et al. [1998]
- 412-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H - 412-D was bound only to B clade virions and to D clade MAL. Nyambi et al. [1998] (subtype comparisons)
- 412-D: Called 412 The tip of the V3 loop was presented in a mucin backbone - higher valency correlates with higher affinity constant. Fontenot et al. [1995] (vaccine antigen design, binding affinity)
- 412-D: Called 412-10D relatively rapid dissociation and weak homologous neutralization. VanCott et al. [1994] (binding affinity)

• 412-D: Neutralizes MN, does not bind SF2 or HXB2 – not reactive with hexa or heptapeptides by Pepscan. Gorny et al. [1993] (variant cross-recognition or cross-neutralization)

• 412-D: Mediated deposition of complement component C3 on HIV infected cells, enhanced by second Ab binding, rabbit anti-human IgG. Spear et al. [1993] (complement)

**No.** 470

MAb ID polyclonal

HXB2 Location gp160 (304-320)

Author Location gp120 (MN)

Epitope RKRIHIGPGRAFYTT

**Neutralizing** L (MN ALA-1)

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Spear et al. 1994

• 40% of antibody in serum that can bind to native viral proteins on MN-infected cells can be blocked by the peptide RKRI-HIGPGRAFYTT, which can also block 75-95% of the complement activation on HIV infected cells. Spear et al. [1994]

**No.** 471

MAb ID CGP 47 439

HXB2 Location gp160 (304-322)

Author Location gp120

Epitope RKRIRIQRGPGRAFVTIGK?

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) human

Ab Type gp120 V3

References Jacobson 1998; Gauduin et al. 1998; Gunthard et al. 1994; Safrit et al. 1993; Liou et al.

- CGP 47 439: Post-exposure passive transfer of murine BAT123 can confer protection to hu-PBL-SCID mice challenged with HIV-1 LAI – this protection is not elicited by CGP 47 439, suggesting that the protection is mediated by complement – the protective ability of BAT123 is lost when mice were treated with cobra venom factor, which inactivates serum complement - in this circumstance complement activation provided a protective advantage. Gauduin et al. [1998]
- CGP 47 439: Review of passive immunotherapy, summarizing Gunthard et al. [1994] in relation to other studies Jacobson [1998]. Gunthard et al. [1994]; Jacobson [1998]
- CGP 47 439: Phase I/IIA clinical trial studying multidose tolerability, immunogenicity and pharmacokinetic responses -GP 47 439 was well tolerated, serum t\_1/2 was 8-16 days, and a virus burden reduction was noted in some patients. Gunthard et al. [1994]
- CGP 47 439: passive transfer to Hu-PBS-SCID mice confers protection against challenge with homologous cell-free virus - CGP 47 439 is a BAT123-human Ig chimera. Safrit et al. [1993]

No. 472

MAb ID polyclonal

**HXB2 Location** gp160 (304–322)

**Author Location (MN)** 

Epitope RKRIHIGPGRAFYTTKN

Subtype multiple

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Cheingsong-Popov et al. 1992

• The Ab response of 829 HIV-1 infected subjects from eight geographic areas to a set of different V3 peptides was determined by ELISA and cross-inhibition studies – the Ab binding pattern was highly variable, depended on the geographic origin of the sample – 297 sera were tested in a neutralization assay – there was a correlation between Ab binding to the MN V3 loop and MN neutralizing titer, but with neutralization of IIIB or CBL-4. Cheingsong-Popov et al. [1992]

No. 473

**MAb ID** 178.1 (178.1.1)

HXB2 Location gp160 (305-309)

**Author Location** gp120 (305–309 BH10)

Epitope KSiRI

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: Env

Species (Isotype) mouse (IgG2a)

Ab Type gp120 V3

Research Contact C. Thiriart, Smith Kline and MRC AIDS rea-

gent project

References Cook et al. 1994; Moore & Ho 1993; Back

et al. 1993; Thiriart et al. 1989

- 178.1: UK Medical Research Council AIDS reagent: ARP331.
- 178.1: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon - this MAb can inhibit gp120 binding to GalCer in vitro - binding of GalCer to gp120 inhibited but did not completely block MAb binding. Cook et al. [1994]
- 178.1: gp41 amino acid substitutions 668 (N/S) and 675 (I/M) in gp41 interfere with 5023s neutralization potency, region 662-675 is ELDKWANLWNWFNI. Back et al. [1993]
- 178.1: Called 178.1.1 conformational, does not bind well to denatured gp120. Moore & Ho [1993]
- 178.1: Reacts to gp120 and gp160 in RIPA EIA and immunoblot. Thiriart et al. [1989]

**No.** 474

MAb ID 257-D (257, 257-2-D-IV, 257-D-IV, 257, 257-2D, 257D, ARP3023)

**HXB2 Location** gp160 (305–309)

Author Location gp120 (MN)

Epitope KRIHI

Subtype B

Neutralizing L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

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References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Zhang et al. 2002; Vella et al. 2002; York et al. 2001; Park et al. 2000; Nyambi et al. 2000; Oggioni et al. 1999; Beddows et al. 1999; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Stamatatos & Cheng-Mayer 1998; Gorny et al. 1998; Yang et al. 1998; LaCasse et al. 1998; Hioe et al. 1997b; Hill et al. 1997; Stamatatos et al. 1997; Schutten et al. 1997; Schutten et al. 1996; Wisnewski et al. 1996; Fontenot et al. 1995; Schutten et al. 1995b; Schutten et al. 1995a; Zolla-Pazner et al. 1995; D'Souza et al. 1995; Stamatatos & Cheng-Mayer 1995; VanCott et al. 1994; D'Souza et al. 1994; Spear et al. 1993; Cavacini et al. 1993a; Gorny et al. 1993; Karwowska et al. 1992b; D'Souza et al. 1991; Gorny et al. 1991

Keywords antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain, assay development, binding affinity, co-receptor, complement, enhancing activity, kinetics, review, subtype comparisons, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-neutralization

- 257-D: UK Medical Research Council AIDS reagent: ARP3023.
- 257-D: NIH AIDS Research and Reference Reagent Program: 1510.
- 257-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 257-D: Called 257: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides do not show as much ability to neutralize as V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. 257 was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 257-D: Called ARP3023: Herpesvirus saimiri-immortalized CD4+ T lymphocytes (HVS T cells) were used to isolate virus and perform HIV-1 neutralization assays, and compared with a standard PBMC protocol neutralization sensitivities to a panel of MAbs and to homologous or heterologous plasma/sera were similar for HVS T cells (CN-2 cells) and PBMCs. Vella *et al.* [2002] (assay development)
- 257-D: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly

sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang *et al.* [2002] (variant cross-recognition or cross-neutralization)

- 257-D: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding. York et al. [2001]
- 257-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding V3 MAbs tended to have the most cross-reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H 257-D showed intermediate reactivity. Nyambi *et al.* [2000] (**subtype comparisons**)
- 257-D: Called 257D six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park *et al.* [2000]
- 257-D: rgp120 derived from a R5X4 subtype B virus, HIV-1 W61D, was used to vaccinate healthy volunteers and the resulting sera were compared with sera from HIV-1 positive subjects and neutralizing MAbs 257-D bound rgp120 W61D but could only neutralize the W61D isolate following T-cell line adaptation. Beddows et al. [1999] (vaccine antigen design, variant cross-recognition or cross-neutralization, vaccine-specific epitope characteristics)
- 257-D: Study of a live-vector mucosal vaccine that expresses HIV-1 V3 domains using the bacterium Streptococcus gordonii which can express heterologous Ag and can colonize the oral cavity and vagina of mice 268-D and 257-D recognized S. gordonii expressing the V3 domain of MN the vaccine stimulated V3-specific IgG2a in mice. Oggioni *et al.* [1999] (vaccine antigen design)
- 257-D: Review of clade specificity and anti-V3 HIV-1-Abs. Zolla-Pazner *et al.* [1999a] (**review**, **subtype comparisons**)
- 257-D: MAb peptide reactivity pattern clustered with immunological related MAbs: 1108, 386, 268, 311, 257, 694.8 the amino acids HI tended to be critical for reactivity in this group. Zolla-Pazner *et al.* [1999b] (antibody binding site definition and exposure)
- 257-D: Kinetic parameters were measured, and the association rates were similar, but dissociation rate constants were quite variable for V3 MAbs, 257-D has a slow dissociation, thus the highest affinity among V3 MAbs. Gorny *et al.* [1998] (kinetics, binding affinity)
- 257-D: A T-cell line-adapted (TCLA) derivative of SI primary isolate 168P acquired the ability to be neutralized by anti-V3 MAbs – the primary isolate could use either CCR5 or CXCR4, and was not neutralized when infection was directed via either pathway, however the TCLA derivative uses CXCR4 only and

is neutralized. LaCasse *et al.* [1998] (**co-receptor**, **variant cross-recognition or cross-neutralization**)

- 257-D: Called 257D deleting the V2 loop of neutralizationresistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F – deletion of V1 or V2 did not enable neutralization by V3 MAbs 391-95D or 257D. Stamatatos & Cheng-Mayer [1998] (vaccine antigen design, subtype comparisons)
- 257-D: A neutralization assay was developed based on heminested PCR amplification of the LTR (HNPCR) LTR-HNPCR consistently revealed HIV DNA and was shown to be a rapid, specific and reliable neutralization assay based on tests with 6 MAbs and 5 isolates. Yang et al. [1998] (assay development)
- 257-D: Called 257 gp120 can inhibit MIP-1alpha from binding to CCR5, but this inhibitory effect is blocked by preincubation of gp120 with three anti-V3 MAbs: 447, 257, 1027 MAb 670 which binds in the C5 region had no effect. Hill et al. [1997] (antibody binding site definition and exposure, co-receptor)
- 257-D: Neutralized (>90%) an SI-env chimeric virus and enhanced (>200%) an NSI-env chimeric virus. Schutten *et al.* [1997] (enhancing activity, variant cross-recognition or cross-neutralization)
- 257-D: Binds less extensively than MAb 391-95D on the surface of HIV-1 isolates SF162 and SF128A neutralizes less potently than 391-95D stronger neutralization of primary macrophage targets than PBMC. Stamatatos *et al.* [1997] (variant cross-recognition or cross-neutralization)
- 257-D: IIIB neutralizing MAbs in vitro fail to neutralize in a mouse model in vivo. Schutten et al. [1996]
- 257-D: 257-D is V H5 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996] (antibody sequence, variable domain)
- 257-D: Called 257-D-IV could neutralize MN and closely related JRCSF, but not 2 B subtype and 1 D subtype primary isolates in a multi-laboratory study involving 11 labs. D'Souza *et al.* [1995] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 257-D: Only inhibition of SI phenotype virus, and strong enhancement of NSI phenotype chimeric viruses, that incorporated different envs from the same donor. Schutten *et al.* [1995a] (enhancing activity, variant cross-recognition or cross-neutralization)
- 257-D: Comparable affinity for SI and NSI viruses, in contrast to MAb MN215. Schutten *et al.* [1995b] (variant cross-recognition or cross-neutralization)
- 257-D: The binding of conformation-dependent anti-V2, anti-V3, and anti-CD4BS MAbs to monomeric and virion-associated gp120 from HIV-1 isolates with differences in cell tropism was studied V3 loop epitopes were less accessible to Ab binding on the virion surface than in the gp120 monomer, particularly for macrophage-tropic isolates SF162 and SF128a, relative to T-cell tropic SF2 sCD4 association with gp120 better revealed this V3 epitope on TCLA SF2 and dual tropic (MU3) viruses than on macrophage tropic isolates. Stamatatos & Cheng-Mayer [1995] (antibody binding

site definition and exposure, variant cross-recognition or cross-neutralization)

- 257-D: In serotyping study using flow-cytometry, bound only to virus with KRIHI. Zolla-Pazner et al. [1995] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 257-D: Included a multi-lab study for antibody characterization and assay comparison – best NAb against MN, but not IIIB.
   D'Souza et al. [1994] (variant cross-recognition or cross-neutralization)
- 257-D: Potent MN neutralization, slow dissociation constant. VanCott *et al.* [1994] (**binding affinity**)
- 257-D: Additive MN or SF2 neutralization when combined with CD4 binding site MAb F105 – does not neutralize RF. Cavacini et al. [1993a] (antibody interactions, variant cross-recognition or cross-neutralization)
- 257-D: Neutralizes MN binds SF2: epitope KSIYI specificity: MN, SF2, NY5, RF. Gorny *et al.* [1993] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 257-D: Mediated deposition of complement component C3 on HIV infected cells, enhanced by second Ab binding, rabbit anti-human IgG – complement mediated virolysis of MN, but not in the presence of sCD4. Spear et al. [1993] (complement)
- 257-D: Reacts with MN, NY5, CDC4 and SF2, does not cross-react with RF, WM52, or HXB2. Karwowska *et al.* [1992b] (variant cross-recognition or cross-neutralization)
- 257-D: Called 257-2-D-IV potent neutralizing MAb. D'Souza et al. [1991]

**No.** 475

**MAb ID** 311-11-D (311-11D, 311, 311D, 311-D)

**HXB2 Location** gp160 (305–313)

Author Location gp120 (MN)

Epitope KRIHIGP

Subtype B

Neutralizing L

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\lambda$ )

Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Gorny et al. 1998; Spear et al. 1993; Gorny et al. 1993; Gorny et al. 1991

**Keywords** antibody binding site definition and exposure, antibody generation, complement, review, subtype comparisons

- 311-11-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 311-11-D: Called 311: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides do not neutralize as

well as V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. 311 was selected using V3 peptides. Gorny *et al.* [2004] (**antibody binding site definition and exposure**)

- 311-11-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding V3 MAbs tended to have the most cross-reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H 311-11-D showed weak reactivity. Nyambi *et al.* [2000] (subtype comparisons)
- 311-11-D: Review of clade specificity and anti-V3 HIV-1-Abs.
   Zolla-Pazner et al. [1999a] (review, subtype comparisons)
- 311-11-D: MAb peptide reactivity pattern clustered with immunological related MAbs: 1108, 386, 268, 311, 257, 694.8

   the amino acids HI tended to be critical for reactivity in this group. Zolla-Pazner *et al.* [1999b] (antibody binding site definition and exposure)
- 311-11-D: Neutralizes MN binds SF2: KSIYIGP. Gorny *et al.* [1993] (antibody binding site definition and exposure, antibody generation)
- 311-11-D: Mediated deposition of complement component C3 on HIV infected cells, enhanced by second Ab binding, rabbit anti-human IgG. Spear *et al.* [1993] (**complement**)

**No.** 476

**MAb ID** 41148D

HXB2 Location gp160 (305-313)

Author Location gp120 (MN)

Epitope KRIHIGP

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 V3

References Gorny & Zolla-Pazner 2004; Alsmadi &

Tilley 1998; Pinter et al. 1993b

**Keywords** ADCC, review, variant cross-recognition or cross-neutralization

- 41148D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. 4117C and 41148D are anti-V3 MAbs that neutralize TCLA strains. Gorny & Zolla-Pazner [2004] (review)
- 41148D: A study of 6 anti-Env MAbs and their ability to bind or direct ADCC against target cells infected with IIIB, MN, SF-2, and RF – bound and directed lysis against strains IIIB, MN, SF-2, comparable to 4117C, however 41148D is 10x less efficient at neutralization, showing ADCC and neutralization don't always correlate. Alsmadi & Tilley [1998] (ADCC)
- 41148D: Neutralizes less potently than 4117C, reacts with MN, IIIB, SF2. Pinter et al. [1993b] (variant cross-recognition or cross-neutralization)

**No.** 477

**MAb ID** 391/95-D (391-95D, 391.5, 391/95D, 391/95)

HXB2 Location gp160 (305–318)

Author Location gp120 (MN)

Epitope KRIHIGPGRAFY

Subtype B

**Neutralizing** L

**Immunogen** HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

References McCaffrey et al. 2004; Gorny et al. 2004;

Gorny & Zolla-Pazner 2004; Zhang et al. 2002; Lawson et al. 2002; Guillon et al. 2002b; Park et al. 2000; Ly & Stamatatos 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Stamatatos & Cheng-Mayer 1998; Stamatatos et al. 1997; Seligman et al. 1996; Stamatatos & Cheng-Mayer 1995; Fontenot et al. 1995; Gorny et al. 1993; Gorny

et al. 1991

**Keywords** acute/early infection, antibody binding site definition and exposure, co-receptor, enhancing activity, review, subtype comparisons, vaccine antigen design, variant cross-recognition

or cross-neutralization

 391/95-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)

- 391/95-D: Called 391/95: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides do not show as much ability to neutralize as V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. 391/95 was selected using V3 peptides. Gorny *et al.* [2004] (antibody binding site definition and exposure)
- 391/95-D: Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The loss of any of the five glycans, within the V3 loop (GM299 V3), C2 (GM292 C2), C3 (GM329 C3), C4 (GM438 C4), or V5 (GM454 V5) made SF162 become sensitive to 391/95-D; SF162 is resistant to 391/95-D neutralization. V3 glycans tended to shield V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans shielded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004] (antibody binding site definition and exposure, vaccine antigen design)
- 391/95-D: The affect of Ab binding on infectivity was studied by pseudotyping three related envs with different phenotypes R5 viruses were preferentially enhanced, not X4 the V3 region was the main determinant of Ab-mediated enhancement and modulation of the interaction between CCR5 and gp120 is critical tests with MAbs anti-V3 391/95-D and CD4BS-specific GP68 indicate that Ab specificity did not determine whether or not infectivity was enhanced or neutralized, rather the phenotype was determined by Env conformation. Guillon et al. [2002b] (co-receptor, enhancing activity)

- 391/95-D: The phenotype and genotype of viral env sequences were studied over a period of seroconversion in one individual Env trans-complementation demonstrated infectivity of clones derived pre-seroconversion were not influenced by MAb 391/95-D, but post-seroconversion clones were enhanced in the presence of 391/95-D, although the V3 binding region was unchanged a change in the CD4-binding site was observed (NL43 427 Glu–>Lys) to be present in the post-seroconversion 391/95-D enhanced clone (see Guillon *et al.* [2002b]) Lawson *et al.* [2002]. Guillon *et al.* [2002b]; Lawson *et al.* [2002] (enhancing activity, acute/early infection)
- 391/95-D: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang *et al.* [2002] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 391/95-D: Called 391-95D SF162 is a neutralization-resistant HIV-1 isolate N-linked glycosylation modifications in the V2 loop of the SF162 gp120 revealed that these sites prevent neutralization by CD4BS MAbs (IgG1b12 and IgGCD4), and protect against neutralization by V3 MAbs (447-D and 391-95D) V2-region glycosylation site mutations did not alter neutralization resistance to V2 MAbs (G3.4 and G3.136) or CD4i MAbs (17b and 48d) V2 glycosylation site modification allows infection of macrophages, probably due to glycosylated forms requiring fewer CCR5 molecules for viral entry. Ly & Stamatatos [2000] (antibody binding site definition and exposure)
- 391/95-D: Called 391/95D six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park *et al.* [2000] (antibody binding site definition and exposure)
- 391/95-D: Review of clade specificity and anti-V3 HIV-1-Abs.
   Zolla-Pazner et al. [1999a] (review, subtype comparisons)
- 391/95-D: Called 391.5 MAb peptide-reactivity pattern clustered with immunological related MAbs: 391.5, 412 and 418, all selected with MN V3 peptide the core amino acids HIGPGR tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure)
- 391/95-D: Called 391-95D deleting the V2 loop of neutralization-resistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F – deletion of V1 or V2 did not enable neutralization by V3 MAbs 391-95D or 257D. Stamatatos &

Cheng-Mayer [1998] (antibody binding site definition and exposure, subtype comparisons)

- 391/95-D: Called 391-95D binds more extensively than MAb 257-D on the surface of HIV-1 isolates SF162 and SF128A neutralizes more potently than 257-D stronger neutralization of primary macrophage targets than PBMC binding postgp120-sCD4 association related to anti-V3 Abs neutralizing capacity. Stamatatos *et al.* [1997] (variant cross-recognition or cross-neutralization)
- 391/95-D: Competition ELISAs with serial deletions estimated the epitope to be KRIHIGPGRAFY – unconstrained peptide had higher affinity than cyclic. Seligman *et al.* [1996] (antibody binding site definition and exposure)
- 391/95-D: The binding of conformation-dependent anti-V2, anti-V3, and anti-CD4BS MAbs to monomeric and virion-associated gp120 from HIV-1 isolates with differences in cell tropism was studied V3 loop epitopes were less accessible to Ab binding on the virion surface than in the gp120 monomer, particularly for macrophage-tropic isolates SF162 and SF128a, relative to T-cell tropic SF2 sCD4 association with gp120 better revealed this V3 epitope on macrophage tropic and dual tropic (MU3) viruses, but not in TCLA SF2. Stamatatos & Cheng-Mayer [1995] (antibody binding site definition and exposure)
- 391/95-D: Neutralizes MN binds to SF2, not IIIB. Gorny et al. [1993]

No. 478

MAb ID Aw

HXB2 Location gp160 (305–320)

**Author Location** gp120 (Gun-1wt)

Epitope KSITIGPGRAFHAI

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: Gun-1 HIV

component: V3

Species (Isotype) rat

Ab Type gp120 V3

References McKnight et al. 1995

Aw: Rat antibodies were raised against V3 peptides that represent either the wildtype (wt), or brain-cell tropic variant (v) of the isolate Gun-1 – Aw gives weak neutralization of both wildtype and v strains. McKnight et al. [1995]

**No.** 479

MAb ID Bw

HXB2 Location gp160 (305–320)

Author Location gp120 (Gun-1wt)

Epitope KSITIGPGRAFHAI

Neutralizing L

Immunogen vaccine

Vector/Type: peptide Strain: Gun-1 HIV

component: V3

Species (Isotype) rat

**Ab Type** gp120 V3

References McKnight et al. 1995

• Bw: Rat antibodies were raised against V3 peptides that represent either the wildtype (wt), or brain-cell tropic variant (v) of the isolate Gun-1 – Bw gives weak neutralization of only the wildtype strain, does not bind to variant. McKnight et al. [1995]

No. 480

MAb ID DO142-10 (DO 142-10)

HXB2 Location gp160 (305–320)

Author Location gp120 (MN)

Epitope KRIHIGPGRAFYTT

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG1) Ab Type gp120 V3

References Gorny & Zolla-Pazner 2004; Kwong et al.

2002; Sullivan et al. 1998a; Parren et al. 1998a; Parren & Burton 1997; Parren et al. 1997b; Ditzel et al. 1997; Seligman et al. 1996

**Keywords** antibody binding site definition and exposure, antibody generation, binding affinity, enhancing activity, review, variant cross-recognition

or cross-neutralization

• DO124-10: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. DO124-10 neutralizes some TCLA strains. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization, review)

- D0124-10: Called D0124. Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-2999) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong et al. [2002] (antibody binding site definition and exposure)
- DO142-10: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 >b3 > b14 > b13 > DO142-10 > DA48 > L17) was markedly different that Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6> DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs – authors suggest that neutralization is determined

by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a] (variant cross-recognition or cross-neutralization, binding affinity)

- DO124-10: The HIV-1 virus YU2 entry can be enhanced by MAbs binding to the CD4BS, V3 loop, and CD4i epitopes – the activation for this enhanced entry state could be conferred on HxB2 by introducing the YU2 V3 loop, or the YU2 V3 and V1/V2 loops – a similar effect is observed by sub-neutralizing concentrations of sCD4 and the effect is dependent of CCR5 – Fab Ab fragment DO124-10 also enhances YU2 entry, ruling out Fc interactions or Env cross-linking as a mechanism - while DO124-10 enhances YU2 entry 6-fold, it neutralizes HXBc2 under identical conditions. Sullivan et al. [1998a] (enhancing activity, variant cross-recognition or cross-neutralization)
- DO142-10: Phage expression libraries panned against MN peptide were used to select Fab DO142-10 - Fab binds MN gp120, but not a primary isolate rec gp120. Ditzel et al. [1997] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- DO142-10: Neutralizes TCLA strains but not primary isolates. Parren et al. [1997b] (variant cross-recognition or cross-neutralization)
- DO142-10: Binds to gp120 MN and an MN V3 peptide with equal affinity, but binds a consensus B peptide and JRCSF less well, and to IIIB gp120 not at all. Parren & Burton [1997] (variant cross-recognition or cross-neutralization, binding affinity)
- DO142-10: Fab fragment competition ELISAs with serial deletions defined the epitope KRIHIGPGRAFYTT. Seligman et al. [1996] (antibody binding site definition and exposure, antibody generation)

**No.** 481

MAb ID Dv

HXB2 Location gp160 (305–320)

Author Location gp120 (Gun-1v)

Epitope KSITIGSGRAFHAI

Neutralizing L

Immunogen vaccine

Vector/Type: peptide Strain: Gun-1 HIV

component: V3

Species (Isotype) rat

Ab Type gp120 V3

References McKnight et al. 1995

• Dv: Rat antibodies were raised against V3 peptides that represent either the wildtype (wt), or brain-cell tropic variant (v) of the isolate Gun-1 – neutralization of only the variant strain, does not bind to wildtype. McKnight et al. [1995]

No. 482

MAb ID Fv

HXB2 Location gp160 (305–320)

Author Location gp120 (Gun-1v)

Epitope KSITIGSGRAFHAI

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: Gun-1 HIV

component: V3

Species (Isotype) rat

Ab Type gp120 V3

References McKnight et al. 1995

• Fy: Rat antibodies were raised against V3 peptides that represent either the wildtype (wt), or brain-cell tropic variant (v) of the isolate Gun-1 – neutralization of only the variant strain, does not bind to wildtype. McKnight et al. [1995]

No. 483

MAb ID Gv

HXB2 Location gp160 (305-320)

Author Location gp120 (Gun-1v)

Epitope KSITIGSGRAFHAI

Neutralizing L

Immunogen vaccine

Vector/Type: peptide Strain: Gun-1 HIV

component: V3

Species (Isotype) rat

Ab Type gp120 V3

References McKnight et al. 1995

• Gv: Rat antibodies were raised against V3 peptides that represent either the wildtype (wt), or brain-cell tropic variant (v) of the isolate Gun-1 - neutralization of only the variant strain, Research Contact Mary White-Scharf, Repligen Corporation, does not bind to wildtype. McKnight et al. [1995]

No. 484

MAb ID Hv

HXB2 Location gp160 (305-320)

Author Location gp120 (Gun-1v)

**Epitope** KSITIGSGRAFHAI

Neutralizing L

Immunogen vaccine

Vector/Type: peptide Strain: Gun-1 HIV

component: V3

Species (Isotype) rat

Ab Type gp120 V3

References McKnight et al. 1995

• Hv: Rat antibodies were raised against V3 peptides that represent either the wildtype (wt), or brain-cell tropic variant (v) of the isolate Gun-1 – neutralization of only the variant strain, does not bind to wildtype. McKnight et al. [1995]

No. 485

MAb ID polyclonal

HXB2 Location gp160 (305-322)

Author Location gp140 (SF162)

Epitope KSITIGPGRAFYATGD

Neutralizing yes

Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade SF162 HIV component:

gp140 Adjuvant: MF59

Species (Isotype) macaque, rabbit (IgG)

Ab Type gp120 V3

References Barnett et al. 2001

• SF162 $\Delta$ V2 is a virus that has a 30 amino acids deletion in the V2 loop that does not abrogate its infectivity but renders it highly susceptible to neutralization – when incorporated into a codon-optimized DNA vaccine with a CMV promoter, delivered by gene gun,  $SF162\Delta2$  gave higher neutralizing Ab titers against SF162 than did SF162 itself, and Abs that crossneutralized non-homologous primary isolates were obtained

only when SF162 $\Delta$ V2, but not intactSF162, was used as the immunogen - NAbs titers specific for SF162 increased with multiple immunizations, while titers for non-homologous isolates decreased, but anti-V3 peptide binding Abs were not likely the source of this distinction because anti-V3 titers were much lower than those against the entire envelope, and the second booster immunization did not increase the titer of anti-V3 loop Abs. Barnett et al. [2001]

No. 486

**MAb ID** 50.1 (R/V3-50.1, Fab 50.1)

HXB2 Location gp160 (306–310)

Author Location gp120 (MN)

**Epitope RIHIG** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN

HIV component: V3

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type gp120 V3

Cambridge, MA

References Zhang et al. 2002; York et al. 2001; Park et al. 2000; Hoffman et al. 1999; Stanfield et al. 1999; LaCasse et al. 1998; Berman

et al. 1997; Seligman et al. 1996; Fontenot et al. 1995; VanCott et al. 1995; Moore et al. 1994b; Robert-Guroff et al. 1994; VanCott et al. 1994; Bou-Habib et al. 1994; Rini et al. 1993; Ghiara et al. 1993; Potts et al. 1993;

White-Scharf et al. 1993; D'Souza et al. 1991

• 50.1: NIH AIDS Research and Reference Reagent Program: 1289

- 50.1: Called R/V3-50.1 A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera - 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 - thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002]
- 50.1: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding - the dissociation constant, Kd of 50.1 for the cell associated primary and TCLA Envs was equal, 7nM. York et al. [2001]
- 50.1: Called R/V3-50.1 six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive - V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the

sensitive form – the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes – 50.1 could only neutralize the sensitive form. Park et al. [2000]

- 50.1: The crystal structure of V3 loop peptides bound to Fabs was obtained – conformational changes in the tip of the V3 loop (GPGR) were observed when different Fabs were bound. Stanfield *et al.* [1999]
- 50.1: A T-cell line-adapted (TCLA) derivative of SI primary isolate 168P acquired the ability to be neutralized by anti-V3 MAbs – the primary isolate could use either CCR5 or CXCR4, and was not neutralized when infection was directed via either pathway, however the TCLA derivative uses CXCR4 only and is neutralized. LaCasse et al. [1998]
- 50.1: Binds to 6/7 isolates from breakthrough cases from a MN gp120 vaccine trial. Berman et al. [1997]
- 50.1: Competition ELISAs with serial deletions produced comparable estimate of epitope length to crystal structure and alanine substitution – KRIHIGP. Seligman et al. [1996]
- 50.1: Used to monitor HIV-1 Env expression in infected H9 cells. VanCott et al. [1995]
- affinity for and neutralization of T cell tropic strain T-CSF, derived from JR-CSF. Bou-Habib et al. [1994]
- 50.1: Shows modest cross-reactivity among B clade gp120s, little outside B clade. Moore et al. [1994b]
- 50.1: Chimeric MN V3 loop in an HXB2 background allows increased FACS signal, Ab affinity, and viral neutralization. Robert-Guroff et al. [1994]
- 50.1: Potent MN neutralization, slow dissociation rate. VanCott et al. [1994]
- 50.1: Crystal structure of a 24 amino acid peptide from the V3 loop bound to 59.1 and 50.1 Fab fragments – epitope KRIHIGP. Ghiara et al. [1993]
- 50.1: No synergistic neutralization of MN when combined with CD4BS MAb F105 - isotype stated to be IgG2a. Potts et al.
- 50.1: Crystal structure of V3 loop bound to 50.1 light chain binds just to the left of GPG, heavy chain binds further to the left. Rini et al. [1993]
- 50.1: Epitope defined by peptide reactivity and changes affinity with amino acid substitutions – epitope RIHIGP. White-Scharf et al. [1993]
- 50.1: Called R/V3-50.1 potent neutralizing of lab strains. D'Souza et al. [1991]

No. 487

MAb ID

HXB2 Location gp160 (306-322)

**Author Location** gp160

**Epitope** RIRPGRAFVTIGK

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: influenza Strain: B clade IIIB

HIV component: V3

Species (Isotype) human (IgA, IgG)

Ab Type gp120 V3

References Garulli et al. 2004

Keywords mucosal immunity

• Progesterone-treated BALB/c mice were intravaginally infected with recombinant influenza A virus (Flu/P18IIIB), expressing the immunodominant CTL epitope (P18IIIB, RIRP-GRAFVTIGK, H-2Dd). A second immunization administered 2 weeks after the first doubled serum IgG levels and enabled detection of vaginal IgG. Low levels of vaginal IgA were detected in some animals. Garulli et al. [2004] (mucosal immunity)

No. 488

MAb ID BAT123 (BAT-123, CGP 47 439)

HXB2 Location gp160 (306–322)

Author Location gp120 (308–322 HXB2)

Epitope RIRIQRGPGRAFVTIGK

Subtype B

**Neutralizing** L

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade IIIB HIV component: HIV-1

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type gp120 V3

• 50.1: No neutralization of primary isolate JR-CSF - greater Research Contact Tanox Biosystems Inc and David Ho, ADARC, NY

References Gauduin et al. 1998; Parren et al. 1998a; Andrus et al. 1998; Poignard et al. 1996a; Sattentau & Moore 1995; Gauduin et al. 1995;

> Pirofski et al. 1993; Thali et al. 1993; Safrit et al. 1993; Moore & Ho 1993; Fung et al. 1990; Liou et al. 1989; Fung et al. 1987

- BAT123: CGP 47 439 is a BAT123 chimera that has a human IgG1 Fc domain.
- BAT123: Post-exposure prophylaxis was effective when MAb 694/98-D was delivered 15 min post-exposure to HIV-1 LAI in hu-PBL-SCID mice, but declined to 50% if delivered 60 min post-exposure, and similar time constraints have been observed for HIVIG, 2F5 and 2G12, in contrast to MAb BAT123 that could protect delivered 4 hours post infection. Andrus et al. [1998]
- BAT123: Post-exposure passive transfer of murine BAT123 can confer protection to hu-PBL-SCID mice challenged with HIV-1 LAI – this protection is not elicited by CGP 47 439, a BAT123 chimera that has a human IgG1 Fc domain, suggesting that the protection is mediated by complement – the protective ability of BAT123 is lost when mice were treated with cobra venom factor, which inactivates serum complement - IgG1 does not fix complement efficiently so an IgG2 MAb might perform better. Gauduin et al. [1998]
- BAT123: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated – authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- BAT123: Epitope described as RGPGRAFVTIGK V3 MAbs 9284, BAT123, 110.5, and 110.I could each significantly increase gp120 dissociation from virus (BAT123 less so than the others), mimicking sCD4, and expose the gp41 epitope for MAb 50-69, in contrast to anti-V2 MAbs. Poignard et al. [1996a]
- BAT123: Passive transfer of BAT123 to hu-PBL-SCID mice 1 hour prior to inoculation with HIV-1 LAI, or up to four hours

post-exposure, could protect mice from infection - the protection, like the MAb, was specific for the viral strain LAI. Gauduin *et al.* [1995]

- BAT123: Binds with high affinity to monomer and oligomer, rapid association and potent neutralization of lab strain. Sattentau & Moore [1995]
- BAT123: Called BAT-123 conformational, does not bind well to denatured gp120 – not reactive with SF-2 gp120 – does not inhibit HIV-1 sera from binding to IIIB gp120. Moore & Ho [1993]
- BAT123: Variable region sequenced heavy chain: V 3660-SB32, D unknown, J H3 – light chain: V kappa21, J kappa2. Pirofski et al. [1993]
- BAT123: Passive transfer to Hu-PBS-SCID mice confers protection against challenge with homologous cell-free virus. Safrit et al. [1993]
- BAT123: Anti-idiotypic MAb, AB19-4i, stimulates anti-anti-ID which neutralizes MN and IIIB. Fung et al. [1990]

No. 489

MAb ID 838-D (838)

HXB2 Location gp160 (307-311)

Author Location Env (RF)

Epitope KSITK

Subtype B

**Neutralizing** L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

Research Contact Susan Zolla-Pazner

(Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

References Gorny et al. 2004; Gorny & Zolla-Pazner

2004; Zhang et al. 2002; He et al. 2002; Nyambi et al. 2000; Gorny et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Nyambi et al. 1998; Hioe et al. 1997b; Gorny et al. 1997

**Keywords** antibody binding site definition and exposure, antibody generation, review, subtype comparisons, variant cross-recognition or crossneutralization

- 838-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- 838-D: Called 838: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 838-D: Called 838 Transgenic mice carrying human genes allowing production of fully human MAbs were used to rapidly create a panel of anti-HIV gp120 MAb producing hybridomas by immunization with HIV SF162 gp120 – the previously described human MAbs 5145A(CD4BS), 4117C (plus others,

V3) and 697D (and SC258, V2) were used as controls. He et al. [2002]

- 838-D: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera – 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 - thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MNprimary strain. Zhang et al. [2002]
- 838-D: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared - no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer - V3 MAbs 447-52D, 838-D, and 1334 bound with a 7-10 fold preference for the oligomer. Gorny et al. [2000] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 838-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H – 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding – V3 MAbs tended to have the most crossreactive binding to clade A, B, C, and D isolates, less to E, F, G, and H – 838-D showed intermediate reactivity. Nyambi et al. [2000] (subtype comparisons)
- 838-D: Review of clade specificity and anti-V3 HIV-1-Abs this Ab showed strong binding to many A, B, C and F peptides, poor binding to D and E. Zolla-Pazner et al. [1999a] (review, subtype comparisons)
- 838-D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 838, 782, 1027, 908, and 1006, all selected with RF V3 peptide - the core amino acids KSITK tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and expo-
- 838-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H – 838-D bound B clade virions but had limited cross-reactivity with other clades, with low levels of binding to A and D virions. Nyambi et al. [1998] (subtype comparisons)
- 838-D: Five human MAbs against were derived from HIVinfected North American subjects after selection by the V3 RF peptide – 838-D was cross-reactive with V3 peptides from clade A and C, and could bind to 5/8 B clade V3 peptides -50% neutralization of RF was obtained. Gorny et al. [1997] (antibody generation, variant cross-recognition or crossneutralization, subtype comparisons)
- 838-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs - BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) - isolates 92HT593 and 91US056

were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations – US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe et al. [1997b] (variant cross-recognition or cross-neutralization)

**No.** 490

MAb ID 1006-15D (1006)

HXB2 Location gp160 (307-312)

Author Location gp120 (RF)

Epitope KSITKG

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zol-Med.

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Center)

References Gorny et al. 2004; Gorny & Zolla-Pazner

2004; He et al. 2002; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. Research Contact Susan

1999a; Gorny et al. 1997

**Keywords** antibody binding site definition and exposure,

antibody generation, review, subtype compar-

- 1006-15D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 1006-15D: Called 1006-15: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 1006-15D: Called 1006 Transgenic mice carrying human genes allowing production of fully human MAbs were used to rapidly create a panel of anti-HIV gp120 MAb producing hybridomas by immunization with HIV SF162 gp120 - the previously described human MAbs 5145A(CD4BS), 4117C (plus others, V3) and 697D (and SC258, V2) were used as controls. He et al. [2002]
- 1006-15D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H – 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding - V3 MAbs tended to have the most crossreactive binding to clade A, B, C, and D isolates, less to E, F, G, and H – 1006-15D showed strong cross-reactivity. Nyambi et al. [2000] (subtype comparisons)
- 1006-15D: Review of clade specificity and anti-V3 HIV-1-Abs - this Ab showed strong binding to several B and F peptides, one C peptide, and some reactivity with A peptides - no binding was observed with D and E peptides. Zolla-Pazner et al. [1999a] (review, subtype comparisons)

• 1006-15D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 838, 782, 1027, 908, and 1006, all selected with RF V3 peptide - the core amino acids KSITK tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure)

1006-15D: Five human MAbs against were derived from HIVinfected North American subjects after selection by the V3 RF peptide – was somewhat cross-reactive with V3 peptides from clade A, C and other B clade V3 peptides, but not E clade. Gorny et al. [1997] (antibody generation, subtype comparisons)

**No.** 491

**MAb ID** 782-D (782)

HXB2 Location gp160 (307–312)

Author Location Env (RF)

Epitope KSITKG

Subtype B

**Neutralizing** L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

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References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Hioe et al. 1997b; Gorny et al. 1997

**Keywords** antibody binding site definition and exposure, antibody generation, review, subtype comparisons, variant cross-recognition or crossneutralization

- 782-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- 782-D: Called 782: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 782-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H - 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding - V3 MAbs tended to have the most crossreactive binding to clade A, B, C, and D isolates, less to E, F, G, and H – 782-D showed intermediate reactivity. Nyambi et al. [2000] (subtype comparisons)
- 782-D: Review of clade specificity and anti-V3 HIV-1-Abs this Ab showed strong binding to several B and F peptides, one C peptide, and some reactivity with A and D peptides. Zolla-Pazner et al. [1999a] (variant cross-recognition or crossneutralization, review, subtype comparisons)

- 782-D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 838, 782, 1027, 908, and 1006, all selected with RF V3 peptide – the core amino acids KSITK tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure)
- 782-D: Five human MAbs against were derived from HIVinfected North American subjects after selection by the V3 RF peptide – 782-D was not cross-reactive with V3 peptides from clade A or E, but could bind to 3/8 B clade V3 peptides, and 1/2 C clade V3 peptides - 50% neutralization of RF was obtained. Gorny et al. [1997] (antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)
- 782-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs - BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) - isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations – US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe et al. [1997b] (variant cross-recognition or cross-neutralization)

No. 492

**MAb ID** 908-D (908, 908-12D)

HXB2 Location gp160 (307-312)

Author Location gp120 (RF)

Epitope KSITKG

Subtype B

**Neutralizing** L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ ) Ab Type gp120 V3

Research Contact Susan Zolla-Pazner

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References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Gorny et al.

**Keywords** antibody binding site definition and exposure, antibody generation, review, subtype comparisons

- 908-D:This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 908-D: Called 908: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was

selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)

- 908-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H - 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding – V3 MAbs tended to have the most cross -reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H – 908-D showed strong cross-reactivity, but achieved only 50% neutralization on 2/5 isolates tested. Nyambi et al. [2000] (subtype comparisons)
- 908-D: Review of clade specificity and anti-V3 HIV-1-Abs - this Ab showed strong binding to several A, B, C and F peptides, and poor binding to E and D peptides. Zolla-Pazner et al. [1999a] (review, subtype comparisons)
- 908-D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 838, 782, 1027, 908, and 1006, all selected with RF V3 peptide - the core amino acids KSITK tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and expo-
- 908-D: Five human MAbs against were derived from HIVinfected North American subjects after selection by the V3 RF peptide - 908-D was not cross-reactive with V3 peptides from clade E, but could bind to 6/8 B clade V3 peptides, 2/4 A clade, and 1/2 C clade - 50% neutralization of RF was obtained. Gorny et al. [1997] (antibody binding site definition and exposure, antibody generation, subtype comparisons)

No. 493

**MAb ID** 1027-15D (1027, 1027-D, 1027D, 1027-15)

HXB2 Location gp160 (307-313)

**Author Location** Env (RF)

Epitope KSITKGP

Subtype B

Neutralizing no

(Zol-

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zol-

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Center)

References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Zhang et al. 2002; Nyambi et al. 2000;

Zolla-Pazner et al. 1999b; Zolla-Pazner et al.

1999a; Gorny et al. 1997

**Keywords** antibody binding site definition and exposure, antibody generation, review, subtype comparisons

- 1027-15D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 1027-15S: Called 1027-15: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)

- 1027-15D: Called 1027-D A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang *et al.* [2002] (antibody binding site definition and exposure)
- 1027-15D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding V3 MAbs tended to have the most cross-reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H 1027-15D showed strong cross-reactivity. Nyambi et al. [2000] (subtype comparisons)
- 1027-15D: Review of clade specificity and anti-V3 HIV-1-Abs

   this Ab showed moderate binding to several B and F peptides,
   one C peptide, and was not reactivity with A, D and E peptides.

   Zolla-Pazner et al. [1999a] (review, subtype comparisons)
- 1027-15D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 838, 782, 1027, 908, and 1006, all selected with RF V3 peptide the core amino acids KSITK tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure)
- 1027-15D: Five human MAbs against were derived from HIV-infected North American subjects after selection by the V3 RF peptide 1027-15D was not cross-reactive with V3 peptides from clade A or E, but could bind to 3/8 B clade V3 peptides, and 1/2 C clade V3 peptides. Gorny *et al.* [1997] (antibody binding site definition and exposure, antibody generation, subtype comparisons)

**No.** 494

MAb ID F19.26-4

**HXB2 Location** gp160 (307–319)

Author Location gp120 (312–324 LAI)

Epitope IRIQRGPGRAFVT

Subtype B

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

**Species** (**Isotype**) mouse ( $IgG2a\kappa$ )

**Ab Type** gp120 V3

References Boudet et al. 1994

F19.26-4: Strain specific – used to raise anti-idiotype antibodies. Boudet et al. [1994]

**No.** 495

**MAb ID** F19.48-3

HXB2 Location gp160 (307-319)

Author Location gp120 (312–324 LAI)

Epitope IRIQRGPGRAFVT

Subtype B

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

**Species** (**Isotype**) mouse ( $IgG2a\kappa$ )

Ab Type gp120 V3

References Boudet et al. 1994

F19.48-3: Strain specific – used to raise anti-idiotype antibodies. Boudet et al. [1994]

**No.** 496

MAb ID F19.57-11

HXB2 Location gp160 (307-319)

Author Location gp120 (312-324 LAI)

Epitope IRIQRGPGRAFVT

Subtype B

**Neutralizing** L (LAI)

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type gp120 V3

**References** Boudet *et al.* 1995; Boudet *et al.* 1994; Boudet *et al.* 1991

- F19.57-11: Anti-anti-idiotypic antibodies (Ab3) were raised in BALBc mice that had greater breadth of reactivity than the original F19.57-11 (Ab3 could also recognize 1282 and SF2, with aa TRK(R or S)IYIGPGRA(WY or FH)T) Boudet et al. [1995]
- F19.57-11: MAb F19.57-11 is strain specific for LAI used to raise anti-idiotype rabbit antibodies (called 57-B Ab2) Boudet *et al.* [1994]

No. 497

**MAb ID** 13105100

HXB2 Location gp160 (307–320)

**Author Location** gp120 (HXB2)

Epitope IRIQRGPGRAFVTI

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

HIV component: V3

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

Research Contact ABI, Columbia, MD

References Dairou et al. 2004

Keywords antibody binding site definition and exposure

• 13105100: This MAb was raised against the peptide IRIQRGP-GRAFVTI, located within the V3 loop flanking the GPGR apical motif. Two MAbs were used to determine the photodamage location in HIV-1 Env induced by sulfonated anionic porphyrins. The negatively charged porphyrins interact with positive charge in the V3 loop. When light activated, they damage amino acid side chains in the C5 region of Env, as evidenced by inhibition of binding of C5 MAb 9201, but not V3 MAb 13105100. Anionic porphyrins could be used in targeted photodynamic decontamination of biological fluids, such as blood, killing HIV without disabling the function of desirable

site definition and exposure)

**No.** 498

MAb ID M77

HXB2 Location gp160 (307-320)

Author Location gp120 (IIIB)

Epitope IRIQRGPGRAFVTI

Neutralizing L

Immunogen HIV-1 infection Species (Isotype) human (IgG) Ab Type gp120 V3

Research Contact Advanced BioScience Laboratories,

Rockville, MD, commercial

References Gorny & Zolla-Pazner 2004; Finnegan et al. 2002; Denisova et al. 2000; Watkins et al. 1996; Denisova et al. 1996; Denisova et al. 1995; DeVico et al. 1995; Cook et al. 1994;

Watkins et al. 1993; di Marzo Veronese et al. 1993; di Marzo Veronese et al. 1992; Pal et al.

1992

Keywords antibody binding site definition and exposure, escape, review, vaccine-specific epitope characteristics, variant cross-recognition or crossneutralization

- M77: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. M77 neutralizes some TCLA strains. Gorny & Zolla-Pazner [2004] (review)
- M77: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I and Cluster II MAbs required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor, binding to a fusion intermediate. Cluster I and II MAbs bound to gp120/gp41 complexes at the cell-to-cell contact interface, in contrast to M77 which bound to gp120 that was evenly dispersed over the target cell surface. Finnegan et al. [2002]
- M77: M77 is highly strain specific for IIIB, but anti-idiotypic Abs directed against M77 can in turn elicit an Ab response with expanded HIV cross-reactivity - this mechanism may serve to prolong the primary response and to counter-balance viral immune evasion by mutation. Denisova et al. [2000] (variant cross-recognition or cross-neutralization)
- M77: Used M77 bound to gp120 as an immunogen analysis of polyclonal and monoclonal (62 MAbs were generated) response suggests the M77-gp120 immunogen generated MAbs to more linear epitopes than gp120 alone or gp120 bound to CD4. Denisova et al. [1996] (vaccine-specific epitope characteristics)
- M77: Native M77 is highly strain specific, and V3 binding is primarily dependent on its heavy chain – a light chain switched Fab version of M77 could recognize HIV-1 strains that had substitutions on the left side of the V3 loop – R in GPGR is likely to be critical for binding. Watkins et al. [1996] (variant cross-recognition or cross-neutralization)
- M77: Reacted with both reduced and non-reduced covalently cross-linked gp120-CD4 complex. DeVico et al. [1995] (antibody binding site definition and exposure)

transfusion products. Dairou et al. [2004] (antibody binding • M77: Conformational rearrangements upon binding of M77 to gp120 generates novel epitopes called metatopes. Denisova et al. [1995]

- M77: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon - this MAb can inhibit gp120 binding to GalCer in vitro. Cook et al. [1994]
- M77: Stated to be a murine MAb a neutralization escape mutant (HXB2 A281V) was selected by growth of HXB2 in the presence of broadly neutralizing sera – M77 neutralization was only slightly reduced by this mutation. Watkins et al. [1993] (escape)
- M77: Antibody binding to viral isolates from IIIB infected lab worker followed through time - A to T substitution resulted in the loss of neutralization and native gp120 binding, but not peptide binding. di Marzo Veronese et al. [1993] (escape)
- M77: IIIB-specific MAb, immunoprecipitates deglycosylated form. di Marzo Veronese et al. [1992] (variant crossrecognition or cross-neutralization)

No. 499

MAb ID polyclonal

HXB2 Location gp160 (307–321)

**Author Location** gp120 (307–321)

Epitope IRIQRGPGRAFVTIG

Subtype B

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) chimpanzee

Ab Type gp120 V3

References Goudsmit et al. 1988

Keywords antibody binding site definition and exposure, autologous responses, variant crossrecognition or cross-neutralization

· By three months post infection, chimpanzees infected with four strains of HIV-1 developed perisistant Ab responses. The V3 loop was a critical binding domain for strain-specific NAbs in sera from the infected chimpanzees. Goudsmit et al. [1988] (antibody binding site definition and exposure, autologous responses, variant cross-recognition or crossneutralization)

No. 500

MAb ID SP.BAL114

**HXB2 Location** gp160 (308–317)

Author Location gp120 (BAL)

**Epitope** SIHIGPGRAF

**Neutralizing** L

**Immunogen** 

**Species (Isotype)** mouse (IgG2a $\kappa$ )

Ab Type gp120 V3

References Arendrup et al. 1995

• Authors suggest that during in vivo immunoselection of escape virus, the V3 domain gains increasing resemblance to that of lab strains. Arendrup et al. [1995]

No. 501

MAb ID SP.SF2:104

HXB2 Location gp160 (308-317)

Author Location gp120 (SF2)

Epitope SIYIGPGRAF

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) (IgG2ak)

Ab Type gp120 V3

References Arendrup et al. 1995; Arendrup et al. 1993

• SP.SF2:104: Authors suggest that during in vivo immunoselection of escape virus, the V3 domain gains increasing resemblance to lab strains. Arendrup et al. [1995]

• SP.SF2:104: Anti-V3 antibody that could neutralize primary virus isolated from a time point of neutralization resistance of autologous virus. Arendrup et al. [1993]

No. 502

MAb ID polyclonal

HXB2 Location gp160 (308–319)

Author Location gp120 (304–318 LAI)

Epitope RIHIGPGRAFYT

Subtype B

Neutralizing

Immunogen HIV-1 infection Species (Isotype) human (IgG, IgM)

Ab Type gp120 V3

References Langedijk et al. 1995

· Polylconal sera from six individuals tested for reactivity against a panel of peptides based on autologous sequences provide evidence for immunological escape mutations in the tip of the V3 loop. Langedijk et al. [1995]

**No.** 503

MAb ID 19b

HXB2 Location gp160 (308-320)

Author Location gp120

Epitope -I----G--FY-T

**Neutralizing** L

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

**Ab Type** gp120 V3

Research Contact James Robinson, University of Connecticut,

References Selvarajah et al. 2005; Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Poignard et al. 2003; Kwong et al. 2002; Zhang et al. 2002; Schulke et al. 2002; Kolchinsky et al. 2001; Park et al. 2000; Binlev et al. 1999; Trkola et al. 1998; Parren et al. 1998a; Mondor et al. 1998; Parren et al. 1997b; Boots et al. 1997; Ugolini et al. 1997; Fouts et al. 1997; Binley et al. 1997a; D'Souza et al. 1997; Trkola et al. 1996a; Wu et al. 1996; Gauduin et al. 1996; Sattentau et al. 1995; Moore & Ho 1995; Moore et al. 1995a; Moore et al. 1995b; Sattentau 1995; Moore et al. 1994a; Moore et al. 1994b; Scott et al. 1990

**Keywords** antibody binding site definition and exposure, antibody interactions, review, vaccine antigen design, vaccine-specific epitope characteristics

• 19b: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. V3 MAbs (447-52D, 19b, F245-B4e8 and 39F) bound to the GDMR antigen, but either did not bind or had diminished binding to mCHO. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)

- 19b: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 19b: This paper attempts to engineer a gp120 molecule that would focus the immune response onto the IgG1b12 epitope. Adding a glycosylation sequon (P313N) to the V3 loop knocked out binding to anti-V3 MAbs loop 2, 19b and 447-52-D. Pantophlet et al. [2003b] (vaccine antigen design)
- 19b: Virion capture assays are not a good predictor of neutralization, and the presentation of epitopes using this assay seems to be different from that of functional Envelope spikes on primary isolates – F105 and b6 could efficiently block the b12-mediated capture of infectious virions in a virus capture, but did not inhibit b12 neutralization – while b12 was potent at neutralizing the three primary virions JR-CSF, A DA, and 89.6, the Abs F105, 19b, and Fab b6 were overall very poor neutralizers. Poignard et al. [2003]
- 19b: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the V3 MAbs used. Zwick et al. [2003] (antibody interactions)
- 19b: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-2999) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face

and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong *et al.* [2002] (antibody binding site definition and exposure)

- 19b: Ab binding characteristics of SOS gp140 were tested using SPR and RIPA SOS gp140 is gp120-gp41 bound by a disulfide bond NAbs 2G12, 2F5, IgG1b12, CD4 inducible 17b, and 19b bound to SOS gp140 better than uncleaved gp140 (gp140unc) and gp120 non-neutralizing MAbs 2.2B (binds to gp41 in gp140unc) and 23A (binds gp120) did not bind SOS gp140. Schulke *et al.* [2002]
- 19b: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002]
- 19b: Mutations in two glycosylation sites in the V2 region of HIV-1 ADA at positions 190 and 197 (187 DNTSYRLINCNTS 199) cause the virus to become CD4-independent and able to enter cells through CCR5 alone – these same mutations tended to increase the neutralization sensitivity of the virus, including to 19b. Kolchinsky et al. [2001]
- 19b: Six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form but 19b was an exception and required around 950 ng/ml to neutralize either form. Park et al. [2000]
- 19b: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits – a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen - SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 - nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999]
- 19b: Used as a control in this Hx10 binding and neutralizing MAb study because 19b does not bind to Hx10. Mondor *et al.* [1998]
- 19b: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated authors sug-

- gest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren *et al.* [1998a]
- 19b: No detectable neutralizing activity among primary isolates with different co-receptor usage – some neutralization of TCLA strains. Trkola et al. [1998]
- 19b: Abs that recognize discontinuous epitopes can identify mimotopes from a phage peptide display library 19b has an epitope involving the tip of the V3 loop, with 5 or 6 essential amino acids distributed within a 12 amino acid stretch the previously determined binding site was confirmed -I—G–FY-T and some tolerated variants described, the I can be I, V, or L, the Y can be Y, F, or W probably a beta-turn is required for FY or FF binding, but WY in can bind with out the context of the turn. Boots *et al.* [1997]
- 19b: In a multilaboratory blinded study, failed to consistently neutralize any of nine B clade primary isolates – there were four sequences with variations in the defined epitope among the 9 isolates tested. D'Souza et al. [1997]
- 19b: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding 19b bound monomer, did not bind oligomer or neutralize JRFL. Fouts et al. [1997]
- 19b: Neutralizes TCLA strains but not primary isolates. Parren *et al.* [1997b]
- 19b: Viral binding inhibition by 19b was weakly correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5) Ugolini *et al.* [1997]
- 19b: Not as effective as IgG1b12 at neutralization *ex vivo* of virus direct from plasma of HIV-1 infected individuals. Gauduin *et al.* [1996]
- 19b: Inhibits gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]
- 19b: MIP-1alpha binding to CCR-5 expressing cells can be inhibited by gp120-sCD4 binding of 19b blocks this inhibition. Wu *et al.* [1996]
- 19b: Binds to some gp120s from clades A,B,C,E, and F weakly neutralized some B and one C clade virus. Moore *et al.* [1995b]
- 19b: Despite broad gp120 binding reactivity, not broadly neutralizing. Moore *et al.* [1995a]
- 19b: Review: more broadly cross-reactive than anti-V3 tip MAb 447-D. Moore & Ho [1995]
- 19b: Formalin inactivation of virus at 0.1% formalin for 10 hours at 4 degrees was optimal for inactivation of virus while maintaining epitope integrity. Sattentau *et al.* [1995]
- 19b: V3 loop binding MAb that is more broadly clade cross-reactive than most (binds to 19/29 clade B and 10/12 clade E gp120s) Moore *et al.* [1994b]
- 19b: Competition studies with human sera from seroconverting individuals showed that anti-CD4 BS antibodies can arise very early in infection, comparable or prior to anti-V3 antibodies. Moore *et al.* [1994a]

No. 504

MAb ID loop 2 (Loop 2, IgG1 Loop 2, loop2)

HXB2 Location gp160 (308–321)

Author Location gp120

Epitope SISGPGRAFYTG

**Neutralizing** L

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

Research Contact D. Burton, Scripps Research Institute, La Jolla, CA

References Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Sullivan et al. 1998a; Parren et al. 1998a; Mondor et al. 1998; Parren & Burton 1997; Parren et al. 1997b; Ugolini et al. 1997; Ditzel et al. 1997; Wu et al. 1996; Moore et al. 1994b; Barbas III et al. 1993

**Keywords** antibody generation, antibody interactions, antibody sequence, variable domain, binding affinity, co-receptor, review, subtype comparisons, vaccine antigen design, variant crossrecognition or cross-neutralization

- loop 2: Also known as Loop 2, IgG1 Loop 2 was a obtained by engineering Fab loop2 into an IgG1 molecule. (antibody generation)
- loop 2: Called loop2. This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. loop 2 neutralizes some TCLA strains. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization, review)
- loop 2: This paper attempts to engineer a gp120 molecule that would focus the immune response onto the IgG1b12 epitope. Adding a glycosylation sequon (P313N) to the V3 loop knocked out binding to anti-V3 MAbs loop 2, 19b and 447-52-D. Pantophlet et al. [2003b] (vaccine antigen design)
- loop 2: Called loop2. scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the V3 MAbs used. Zwick et al. [2003] (antibody interactions)
- loop 2: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13> DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs - authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope - binding affinity of divalent IgG1 loop 2 is only 2-fold greater than monovalent Fab loop 2, suggesting the IgG1 form may bind with only one arm. Parren et al. [1998a] (binding affinity)
- loop 2: The HIV-1 virus YU2 entry can be enhanced by MAbs binding to the CD4BS, V3 loop, and CD4i epitopes - the activation for this enhanced entry state could be conferred on HxB2 by introducing the YU2 V3 loop, or the YU2 V3 and

V1/V2 loops – a similar effect is observed by sub-neutralizing concentrations of sCD4 and the effect is dependent of CCR5 loop 2 enhances YU2 at concentrations up to 20 ug/ml. Sullivan et al. [1998a]

- loop 2: Binds to gp120 from MN and SF2 but not LAI. Ditzel et al. [1997] (variant cross-recognition or crossneutralization)
- loop 2: Epitope is suggested to be GPGRAF binds to 10/17 US clade B monomeric gp120s – IgG1 form can neutralize MN and 2 primary isolates tested. Parren & Burton [1997]
- loop 2: Neutralizes TCLA strains but not primary isolates. Parren et al. [1997b] (variant cross-recognition or crossneutralization)
- loop 2: Viral binding inhibition by loop 2 MAb or Fab was correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5) Ugolini et al. [1997]
- loop 2: MIP-1alpha binding to CCR-5 expressing cells can be inhibited by gp120-sCD4 – binding of loop 2 blocks this inhibition. Wu et al. [1996] (co-receptor)
- loop 2: Called Loop 2 shows modest cross-reactivity among B clade gp120s, little outside B clade. Moore et al. [1994b] (variant cross-recognition or cross-neutralization, subtype comparisons)
- loop 2: Sequences of the heavy and light chain Fab variable regions were generated. Barbas III et al. [1993] (antibody sequence, variable domain)

No. 505

**MAb ID** 4G10

HXB2 Location gp160 (308-322)

Author Location gp120 (308–322 LAI)

Epitope RIQRGPGRAFVTGK

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: HBcAg fusion HIV compo-

nent: V3

Species (Isotype) mouse

Ab Type gp120 V3

Research Contact Dr.

Albrecht von Brunn, Max-von-Pettenkofer-Institut, Ludwig-Maximilians-

Universitat Munchen, Germany

References von Brunn et al. 1993

- 4G10: NIH AIDS Research and Reference Reagent Program:
- 4G10: A 25 amino acid V3-loop sequence fused to HBcAg enhanced V3 immunogenicity. von Brunn et al. [1993]

No. 506

MAb ID 5F7

HXB2 Location gp160 (308–322)

Author Location gp120 (308–322 LAI)

Epitope RIQRGPGRAFVTGK

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: HBcAg fusion HIV compo-

nent: V3

Species (Isotype) mouse

Ab Type gp120 V3

Research Contact Dr. Albrecht von Brunn, Max-von- Author Location gp120 (IIIB) Pettenkofer-Institut, Ludwig-Maximilians-

Universitat Munchen, Germany

References von Brunn et al. 1993

• 5F7: NIH AIDS Research and Reference Reagent Program: Species (Isotype) mouse 2533.

• 5F7: A 25 amino acid V3-loop sequence fused to HBcAg Research Contact Dupont, commercial enhanced V3 immunogenicity. von Brunn et al. [1993]

No. 507

**MAb ID** G3-523

HXB2 Location gp160 (308-322) Author Location gp120 (308-322) Epitope RIQRGPGRAFVTIGK

> **Neutralizing Immunogen**

Species (Isotype) mouse

Ab Type gp120 V3

References Jagodzinski et al. 1996; Matsushita et al. 1988

• G3-523: The sulfated polysaccharide curdlan sulfate (CRDS) Research Contact Abraham Pinter, Public Health Research Inbinds to the Envelope of T-tropic viruses and neutralizes virus – CRDS inhibits G3-523 binding. Jagodzinski et al. [1996]

No. 508

MAb ID MN215

HXB2 Location gp160 (308-322)

Author Location gp120 (MN)

**Epitope RIHIGPGRAFYTTKN** 

**Neutralizing** L

**Immunogen** HIV-1 infection Species (Isotype) human (IgG1)

Ab Type gp120 V3

References Zipeto et al. 2005; Gorny & Zolla-Pazner 2004; Schutten et al. 1995b

**Keywords** antibody binding site definition and exposure, review, vaccine antigen design, variant crossrecognition or cross-neutralization

- MN215: MRC Centralized Facility for AIDS Reagents, NIBSC, UK, EVA3056
- MN215: HIV-1 fusion complexes were prepared from cell lines expressing R5 HIV-1 gp120/gp41 and CD4-CCR5. Neutralizing Abs were raised against both R5 (strain BaL) and X4 (strain 213) viruses. MN215 was used to detect gp120/gp41. Zipeto et al. [2005] (vaccine antigen design)
- MN215: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. MN215 neutralizes some TCLA strains. Gorny & Zolla-Pazner [2004] (review)
- MN215: Minimum epitope for MAB using the Dutch consensus is AFYTTGE, different than defined for MN - generated by EBV transformation of PBMC - displayed higher affinity for NSI than for SI glycoproteins - amino acids HIGP were essential for binding. Schutten et al. [1995b] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)

No. 509

MAb ID Nea 9301

HXB2 Location gp160 (308-323)

Epitope RIQRGPGRAFVTIGKI

Neutralizing

**Immunogen** 

Ab Type gp120 V3

References Wagner et al. 1996

**No.** 510

**MAb ID** 4117C (4117c)

HXB2 Location gp160 (309-315)

Author Location gp120

Epitope IXIGPGR

Subtype B

Neutralizing L

Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\lambda$ )

Ab Type gp120 V3

stitute, Newark, NJ, 07103. pinter@phri.org.

References Pinter et al. 2005; Krachmarov et al. 2005; Pinter et al. 2004; Gorny & Zolla-Pazner 2004; He et al. 2002; Alsmadi & Tilley 1998; Pinter et al. 1993b; Pinter et al. 1993a;

di Marzo Veronese et al. 1993; Tilley et al.

1992; Tilley et al. 1991a

Keywords ADCC, antibody binding site definition and exposure, antibody interactions, review, subtype comparisons, variant cross-recognition

or cross-neutralization

- 4117c: Sera from subtype A infected individuals from Cameroon have antibodies that react strongly with subtype A and subtype B V3 loops in fusion proteins, and neutralize SF162 pseudotypes, while sera from subtype B infected individuals reacted only with subtype B. Sera from Cameroon did not neutralize primary A or B isolates, due to indirect masking by the V1/V2 domain rather than due to loss of the target epitope. Neutralization by anti-V3 B clade specific MAbs 447-52D and 4117C was fully blocked by a clade V3 loop fusion protein, but not an A clade fusion protein, while Cameroonian sera neutralization was fully blocked by both A and B clade fusion proteins. Krachmarov et al. [2005] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 4117c: This study is about the MAb C108g, and 4117C was a control. 4117C is a linear V3 epitope unaffected by reduction, whereas C108g, contrary to earlier reports, requires disulfide bonds. C108G is type-specific and neutralizes BaL and HXB2. JR-FL is a neutralization resistant strain; modification of JRFL at V2 positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The modification in V2 also increased neutralization sensitivity to V3 MABs 4117c, 2219, 2191, and 447-52D, but only had minor effects on neutralization by CD4BS MAb 5145A, and broadly neutralizing MAbs IgG1b12, 2G12, and 2F5. Binding to CCR5 was completely inhibited by two V3 MAbs, 4117C and 2219, and was substantially inhibited by 2G12, but was not inhibited by C108g. Pinter et al. [2005] (antibody binding site definition and exposure)

- 4117c: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. 4117C and 4118D are anti-V3 MAbs that neutralize TCLA strains. Gorny & Zolla-Pazner [2004] (review)
- 4117c: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12 which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. 5/6 anti-V3 MAbs, including 4117c, had similar binding affinity to soluble SF162 and JR-FL rgp120s, although the V3 loop differs at three positions (HigpgrafyTtgE for JR-FL and TigpgrafyAtgD for SF162). Pinter et al. [2004] (variant cross-recognition or cross-neutralization)
- 4117C: Transgenic mice carrying human genes allowing production of fully human MAbs were used to rapidly create a panel of anti-HIV gp120 MAb producing hybridomas by immunization with HIV SF162 gp120 the previously described human MAbs 5145A(CD4BS), 4117C (plus others, V3) and 697D (and SC258, V2) were used as controls. He *et al.* [2002]
- 4117C: A study of 6 anti-Env MAbs and their ability to bind or direct ADCC against target cells infected with IIIB, MN, SF-2, and RF – bound and directed lysis against MN and SF2, but not IIIB and RF. Alsmadi & Tilley [1998] (ADCC, variant cross-recognition or cross-neutralization)
- 4117C: Neutralizes SF2 and MN synergistically combined with anti-CD4 binding site discontinuous MAb. Pinter *et al.* [1993a]; Tilley *et al.* [1992] (antibody interactions, variant cross-recognition or cross-neutralization)
- 4117C: Binds V3 loop does not immunoprecipitate soluble gp120, does react with gp120 on intact virions. Pinter *et al.* [1993b] (antibody binding site definition and exposure)
- 4117C: Potent neutralizing activity against MN, SF-2, and NY-5 – synergy with CD4BS MAb 1125H. Tilley et al. [1991a] (antibody binding site definition and exposure, antibody interactions, variant cross-recognition or cross-neutralization)

**No.** 511

**MAb ID** 419-D (419, 419D)

HXB2 Location gp160 (309-315)

**Author Location** gp120 (MN)

**Epitope IHIGPGR** 

Subtype B

Neutralizing L

Immunogen HIV-1 infection

**Species (Isotype)** human ( $IgG1\lambda$ )

Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

**References** Gorny *et al.* 2004; Gorny & Zolla-Pazner 2004; He *et al.* 2002; Nyambi *et al.* 2000; Zolla-Pazner *et al.* 1999b; Zolla-Pazner *et al.* 1999a; Nyambi *et al.* 1998; Hioe *et al.* 1997b;

Fontenot *et al.* 1995; Spear *et al.* 1993; Gorny *et al.* 1993; Karwowska *et al.* 1992b

**Keywords** antibody binding site definition and exposure, complement, mimotopes, review, subtype comparisons, superinfection, variant cross-recognition or cross-neutralization

- 419-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 419-D: Called 419: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (mimotopes, superinfection)
- 419-D: Called 419 Transgenic mice carrying human genes allowing production of fully human MAbs were used to rapidly create a panel of anti-HIV gp120 MAb producing hybridomas by immunization with HIV SF162 gp120 – the previously described human MAbs 5145A(CD4BS), 4117C (plus others, V3) and 697D (and SC258, V2) were used as controls. He *et al.* [2002]
- 419-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding V3 MAbs tended to have the most cross-reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H 419-D showed intermediate reactivity, and no neutralization when tested against five strains discrepancy between the epitope as described in earlier papers and as described here, KRIHIGP. Nyambi *et al.* [2000] (**subtype comparisons**)
- 419-D: Review of clade specificity and anti-V3 HIV-1-Abs epitope is described as KRIHIGP. Zolla-Pazner *et al.* [1999a] (antibody binding site definition and exposure, review)
- 419-D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 1334, 419, 504, 447, 453 and 537 the core amino acids GP tended to be critical for reactivity in this group. Zolla-Pazner *et al.* [1999b] (antibody binding site definition and exposure)
- 419-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H 419-D bound to 3/4 B clade virions, and to D clade MAL. Nyambi *et al.* [1998] (**subtype comparisons**)
- 419-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D,

450-D, 670-D, 1281-D and 98-6. Hioe et al. [1997b] (variant cross-recognition or cross-neutralization)

- 419-D: Neutralizes MN binds SF2: IYIGPGR. Gorny et al. [1993] (variant cross-recognition or cross-neutralization)
- 419-D: Mediated deposition of complement component C3 on HIV infected cells, enhanced by second Ab binding, rabbit anti-human IgG. Spear et al. [1993] (complement)
- 419-D: MN, NY5 and SF2 strain specific, does not cross-react with RF, CDC4, WM52 or HXB2. Karwowska et al. [1992b] (variant cross-recognition or cross-neutralization)

No. 512

**MAb ID** 453-D (453)

HXB2 Location gp160 (309-315)

Author Location gp120 (MN)

Epitope IHIGPGR

Subtype B

Neutralizing L

Immunogen HIV-1 infection **Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

Research Contact Susan

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Center)

References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Fontenot et al. 1995; VanCott et al. 1994; Gorny et al. 1993; Gorny et al. 1991

Keywords antibody binding site definition and exposure, binding affinity, review, subtype comparisons, vaccine antigen design, variant crossrecognition or cross-neutralization

- 453-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization)
- 453-D: Called 453: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 453-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H - 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding – V3 MAbs tended to have the most crossreactive binding to clade A, B, C, and D isolates, less to E, F, G, and H – 453-D showed intermediate reactivity. Nyambi et al. [2000] (subtype comparisons)
- 453-D: Review of clade specificity and anti-V3 HIV-1-Abs. 504-D: MAb peptide-reactivity pattern clustered with immuno-Zolla-Pazner *et al.* [1999a] (**review**, **subtype comparisons**)
- 453-D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 1334, 419, 504, 447, 453 and 537 - the core amino acids GP tended to be critical for reactivity in this group – MAb 268, with a previously defined core epitope identical to 453 (HIGPGR), was not part of this reactivity group,

illustrating that context can be critical. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure)

- 453-D: Called 453, epitope described as KRIHIGPGR the tip of the V3 loop was presented in a mucin backbone - higher valency correlates with stronger affinity constant. Fontenot et al. [1995] (antibody binding site definition and exposure, vaccine antigen design)
- 453-D: Moderate homologous neutralization, moderately slow dissociation rate. VanCott et al. [1994] (binding affinity)
- 453-D: Neutralizes MN binds SF2: IYIGPGR specificity: MN, SF2, NY5, RF. Gorny et al. [1993] (antibody binding site definition and exposure)

**No.** 513

**MAb ID** 504-D (504, 504-10D)

HXB2 Location gp160 (309–315)

Author Location gp120 (MN)

Epitope IHIGPGR

Subtype B

**Neutralizing** L

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

> References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Gorny et al. 1993

> **Keywords** antibody binding site definition and exposure, review, subtype comparisons

- 504-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 504-D: Called 504: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 504-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H – 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding - V3 MAbs tended to have the most crossreactive binding to clade A, B, C, and D isolates, less to E, F, G, and H – 504-D showed weak reactivity. Nyambi et al. [2000] (subtype comparisons)
- 504-D: Review of clade specificity and anti-V3 HIV-1-Abs. Zolla-Pazner *et al.* [1999a] (**review**)
- logical related MAbs: 1334, 419, 504, 447, 453 and 537 the core amino acids GP tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure)
- 504-D Neutralizes MN binds SF2: IYIGPGR. Gorny et al. [1993] (antibody binding site definition and exposure)

No. 514

**MAb ID** 83.1 (MAb 83.1)

HXB2 Location gp160 (309-315)

Author Location gp120 (SF2)

Epitope IYIGPGR

Neutralizing L

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN

HIV component: V3

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

Research Contact Mary White-Scharf, Repligen Corporation,

Cambridge, MA

References Binley et al. 1999; Keller & Arora 1999;

Jelonek et al. 1999; Potts et al. 1993; White-

Scharf et al. 1993

- 83.1: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits – a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen – SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 – nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 - MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999]
- 83.1: Maternally transferred anti-V3 loop MAb selectively inhibits the anti-V3 loop Ab component of the IgG response to rgp120 SF2 in 21 day old BALBc mice. Jelonek et al. [1999]
- 83.1: 19 day old mice injected with 83.1 have a shift in IgG1 response away from the V3 loop upon vaccination, without decreasing the total IgG anti-gp120 response, suggesting that prior treatment with a MAb can mask immunogenic sites and shift the immune response to vaccination. Keller & Arora [1999]
- 83.1: Study of synergism of neutralization and binding comparing F105 and sCD4 with the V3 MAbs: 50.1, 59.1, 83.1, and 58.2 – synergy was observed, and the data suggest that binding of one ligand (F105) can increase the binding of the second (e. g. V3 loop MAbs) due to conformational changes. Potts et al.
- 83.1: Neutralizes SF2. White-Scharf et al. [1993]

No. 515

**MAb ID** 5023B

**HXB2 Location** gp160 (309–316)

**Author Location** gp120 (309–316 BH10)

Epitope IQRGPGra

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade BH10 Author Location gp120

HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Langedijk et al. 1991

• 5023B: Generation and fine mapping of murine MAbs. Langedijk et al. [1991]

**No.** 516

MAb ID F58/D1 (F58)

HXB2 Location gp160 (309–316)

Author Location gp120 (IIIB)

Epitope IxxGPGRA

Neutralizing L

Immunogen vaccine

Vector/Type: virus derived protein HIV com-

ponent: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

References Heap et al. 2005b; Jackson et al. 1999; Millar

et al. 1998; Moore et al. 1993b; Levi et al. 1993; Broliden et al. 1991; Akerblom et al.

1990

Keywords antibody binding site definition and exposure, antibody sequence, variable domain, structure

- F58/D1: Called F58. A 17 amino acid peptide from the CDR-H3 region of F58 retained specificity for gp120 and could neutralize IIIB, although less efficiently than the intact antibody. The F58 MicroAb has a 3-fold faster association rate and a 37.5fold more rapid dissociation rate than the intact antibody. Such Ab binding site fragments, that retain binding specificity, are called microantibodies. Alanine-substitutions in F58 MicroAb at three positions significantly compromised neutralization but did not reduce binding to soluble gp120, while substitutions at three other positions abrogated both binding and neutralization. The microAb forms a conformationally constrained beta sheet. Heap et al. [2005b] (antibody binding site definition and exposure, antibody sequence, variable domain, structure)
- F58/D1: A 17 amino acid MicroAB was made from the third complementarity-determining region of the heavy chain of MAb – F58 neutralized 5x's more efficiently in terms of mass than the original MAb, 32-fold less on a molar basis – neutralization does not involve initial attachment, but fusion and events in early infection. Jackson et al. [1999]
- F58/D1: The interaction of a 17-amino-acid neutralizing microantibody (MicroAB) based on F58 and HIV-1 env was studied by electrospray ionization mass spectrometry. Millar et al. [1998]
- F58/D1: Called F58. The complementarity-determining region of F58 was used to create a miniantibody that could neutralize both HIV-1 IIIB and SF2 in vitro. Levi et al. [1993] (antibody binding site definition and exposure, antibody sequence, variable domain)
- F58/D1: Binding to native gp120 1-3 fold greater than to denatured – 314G/W substitution abolishes binding, changes outside the loop have little effect. Moore et al. [1993b]

No. 517

MAb ID P1/D12

**HXB2 Location** gp160 (309–316)

Epitope IxxGPGRA

Neutralizing L

Immunogen vaccine

Vector/Type: virus derived protein Strain: B

clade IIIB HIV component: gp120 Epitope IQRGPGRAF

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Moore et al. 1993b; Akerblom et al. 1990

• P1/D12: Binding to native gp120 1-3 fold greater than to denatured – 314G/W substitution abolishes binding, changes outside the loop have little effect. Moore et al. [1993b]

**No.** 518

**MAb ID** P4/D10 (P4D10)

HXB2 Location gp160 (309-316)

Author Location gp120

Epitope IxxGPGRA

Neutralizing L

Immunogen vaccine

clade IIIB HIV component: gp120

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type gp120 V3

References Schonning et al. 1999; Schonning et al. 1998; Jacobson 1998; Hinkula et al. 1994; Arendrup et al. 1993; Moore et al. 1993b; Marks et al. 1992; Broliden et al. 1991; Broliden et al. 1990; Akerblom et al. 1990

- P4/D10: Called P4D10 the stoichiometry of MAb neutralization was tested and the data indicated that binding for neutralization was was incremental not all or none, i.e., each envelope oligomer binds a single MAb and each Env oligomer bound reduces the chances of infection - MAb BC1071 was used for virion quantification - P4D10 binds only to Env with a glycosylation site mutation at the base of the V3 loop A308T. Schonning et al. [1999]
- P4/D10: Review of passive immunotherapy, summarizing Hinkula et al. [1994] in relation to other studies Jacobson [1998]. Hinkula et al. [1994]; Jacobson [1998]
- P4/D10: Called P4D10 In a study of the influence of the glycan at position 306 of the V3 loop on MAb recognition, anti-V3 MAbs were found to neutralize an HIV-BRU mutant virus that lacks the V3 loop glycan more efficiently than HIV-BRU – Ab binding site was suggested to be 314-323 of BRU. Schonning et al. [1998]
- P4/D10: Used for passive immunotherapy in four late-stage HIV-infected patients – the serum level of p24 did not decrease in any of these four - see also MAb F58/H3. Hinkula et al. [1994]
- P4/D10: Primary isolates from different time points from one individual were not susceptible to neutralization by P4/D10. Arendrup et al. [1993]
- P4/D10: Binding to native gp120 3 fold greater than to denatured – 314G/W substitution abolishes binding, changes outside the loop have little effect. Moore et al. [1993b]
- P4/D10: Variable domain sequenced and is identical to F58/H3. Marks et al. [1992]
- P4/D10: Neutralizing and ADCC activity. Broliden et al. [1990]

No. 519

**MAb ID** IIIB-13 V3 (1044-13 IIIB-V3-13 1727)

HXB2 Location gp160 (309–317)

Author Location gp120 (308–316 IIIB)

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

References Zhang et al. 2002; Chakrabarti et al. 2002; Watkins et al. 1993; D'Souza et al. 1994; Laman et al. 1993; Laman et al. 1992

- IIIB-13 V3: Also known as 1044-13 and as IIIB-V3-13 (J. P. Moore, per. comm.)
- IIIB-13 V3: UK Medical Research Council AIDS reagent: ARP3046.
- IIIB-13 V3: NIH AIDS Research and Reference Reagent Program: 1727.
- Vector/Type: virus derived protein Strain: B IIIB-13 V3: Called 1727: Used as a standard for comparing immune responses to modified gp140 (gp140deltaCFI), with C-term mutations intended to mimic a fusion intermediate and stabilize trimer formation - experiment showed enhanced humoral immunity without diminishing the CTL response in mice injected with a DNA vaccine. Chakrabarti et al. [2002]
  - IIIB-13 V3: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera - 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 – thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002]
  - IIIB-13 V3: Included in a panel of antibodies used in a multilab study for antibody characterization and assay comparison, some neutralization of strains other than IIIB. D'Souza et al. [1994]
  - IIIB-13 V3: Called IIIB-V3-13 a neutralization escape mutant (HXB2 A281V) was selected by growth of HXB2 in the presence of broadly neutralizing sera - IIIB-V3-13 neutralization was only slightly reduced by this mutation. Watkins et al. [1993]
  - IIIB-13 V3: Neutralizes IIIB but not MN. Laman et al. [1992]

No. 520

**MAb ID** IIIB-34 V3 (IIIB-V3-34)

**HXB2 Location** gp160 (309–317)

Author Location gp120 (308–316 IIIB)

**Epitope IQRGPGRAF** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

**Species (Isotype)** mouse (IgG1)

Ab Type gp120 V3

References Laman et al. 1993; Laman et al. 1992

• IIIB-34 V3: UK Medical Research Council AIDS reagent: ARP3047.

ization – binding is reduced somewhat by DTT or SDS-DTT. enhanced by NP40, but binds to native and denatured gp120. Laman et al. [1993]

• IIIB-34 V3: Neutralizes IIIB but not MN – QXGPG are critical amino acids for binding by Pepscan analysis. Laman et al. [1992]

No. 521

**MAb ID** A47/B1

HXB2 Location gp160 (309–318)

Author Location gp120 (307–316 IIIB)

Epitope IQRGPGRAFV

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Akerblom et al. 1990

No. 522

MAb ID D59/A2

**HXB2 Location** gp160 (309–318)

Author Location gp120 (307–316 IIIB)

Epitope IQRGPGRAFV

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Akerblom et al. 1990

No. 523

MAb ID G44/H7

HXB2 Location gp160 (309-318)

**Author Location** gp120 (307–316 IIIB)

Epitope IQRGPGRAFV

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Akerblom et al. 1990

No. 524

MAb ID MO96/V3 (M096, M096/V3)

HXB2 Location gp160 (309-318)

Author Location gp120 (309-318)

Epitope IQRGPGRAFV+AHCNISRAKW

Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgM)

Ab Type gp120 V3

References Gorny & Zolla-Pazner 2004; Ohlin et al. 1992

Keywords antibody binding site definition and exposure,

antibody generation, review

• IIIB-34 V3: Called IIIB-V3-34 – IIIB strain specific neutral- • M093/V3: Review, provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains; a subset can also neutralize some primary isolates. The three IgMs, M096, M097, and M099, are nonneutralizing. Gorny & Zolla-Pazner [2004] (review)

> M096/V3: Generated in response to IIIB Env 286-467 upon in vitro stimulation of uninfected-donor lymphocytes, and binds to two peptides: 309-318 + 329-338. Ohlin et al. [1992] (antibody binding site definition and exposure, antibody generation)

> > No. 525

**MAb ID**  $\mu$ 5.5 (5.5, mu5.5, Rmu5.5)

HXB2 Location gp160 (309-319)

Author Location gp120 (MN)

Epitope IHIGPGRAFYT

**Neutralizing** LP

Immunogen

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type gp120 V3

References Okamoto et al. 1998; Maeda et al. 1992

- mu5.5: Rmu5.5 is a humanized antibody of mouse MAb m5.5 - neutralized primary isolates with similar V3 loops - passive transfer of MAb to SCID-hu or hu-PBL-SCID mice conferred protection. Okamoto et al. [1998]
- mu5.5: sCD4 causes loss of IIIB type-specificity for MAb 0.5beta, allowing binding and neutralization of MN, in contrast to MAb mu5.5. Maeda et al. [1992]

No. 526

MAb ID 268-D (268-11-D-IV, 268D, 268, 268-11D, 268-10D, MAb 268, 268-10-D, ARP)

HXB2 Location gp160 (310–315)

Author Location gp120 (MN)

Epitope HIGPGR

Subtype B

**Neutralizing** L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

(Zol-Research Contact Susan Zolla-Pazner

> las01@mcrcr6.med.nyu) (NYU Med.

Center)

References Lusso et al. 2005; Gorny et al. 2004; Gorny &

Zolla-Pazner 2004; Zhang et al. 2002; Vella et al. 2002; York et al. 2001; Park et al. 2000; Nyambi et al. 2000; Hioe et al. 2000; Laisney & Strosberg 1999; Oggioni et al. 1999; Beddows et al. 1999; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; LaCasse et al. 1998; Stamatatos et al. 1997; Hioe et al. 1997b; Wisnewski et al. 1996; McKeating et al. 1996; Fontenot et al. 1995; Zolla-Pazner et al. 1995; Stamatatos & Cheng-Mayer 1995; VanCott et al. 1994; Spear et al. 1993; Gorny et al. 1993; Karwowska et al. 1992b; D'Souza

et al. 1991; Gorny et al. 1991 **Keywords** antibody binding site definition and exposure,

review

- 268-D: UK Medical Research Council AIDS reagent:
   268-D: A panel of 47 human MAbs was tested against 26 HIV-1 ARP3024.
   group M primary isolates from clades A through H – 19 V3
- 268-D: NIH AIDS Research and Reference Reagent Program: 1511.
- 268-D: The epitope for the MAb D19 is conserved and embedded in V3. D19 is unique in that for R5 viruses, it was cryptic and did not bind without exposure to sCD4, and for X4 and R5X4 isolates it was constitutively exposed. It had an overlapping binding region with MAbs 447-52D, B4e8, and 268-D, but different reactivity patterns and fine specificity. Lusso *et al.* [2005]
- 268-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, while many neutralize some TCLA strains, a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 268-D: Called 268: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4 induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides do not show as much ability to neutralize as V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. 268 was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 268-D: Called ARP3024: Herpesvirus saimiri-immortalized CD4+ T lymphocytes (HVS T cells) were used to isolate virus and perform HIV-1 neutralization assays, and compared with a standard PBMC protocol – neutralization sensitivities to a panel of MAbs and to homologous or heterologous plasma/sera were similar for HVS T cells (CN-2 cells) and PBMCs. Vella et al. [2002]
- 268-D: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera—2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5—thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002]
- 268-D: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding one of the TCLA V3 viruses 320SI-C3.3 shows reduced binding with this MAb, the sequence of the epitope in 320SI is HIGPGR and in 320SI-C3.3 is RIGPGR. York et al. [2001]
- 268-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses CD4BS MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells V3 MAbs 447-52-D and 268-10-D did not effect proliferation. Hioe *et al.* [2000]

- 268-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding V3 MAbs tended to have the most cross-reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H 268-D showed weak reactivity. Nyambi *et al.* [2000]
- 268-D: Called 268D six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park et al. [2000]
- 268-D: rgp120 derived from a R5X4 subtype B virus, HIV-1 W61D, was used to vaccinate healthy volunteers and the resulting sera were compared with sera from HIV-1 positive subjects and neutralizing MAbs 268-D bound rgp120 W61D but could only neutralize the W61D isolate following T-cell line adaptation. Beddows *et al.* [1999]
- 268-D: Called MAb 268 To identify potential mimotopes of V3, a hexapeptide phage library was screened with MAb 268 – two hexamers were identified, HLGPGR or KAIHRI that bind to 268 with the same binding site as the V3 loop and inhibit 268 MN gp120 – KLH conjugated hexamer KAIHRI stimulates Abs in rabbits that cross-react with ML gp120. Laisney & Strosberg [1999]
- 268-D: Called 268-11D Study of a live-vector mucosal vaccine that expresses HIV-1 V3 domains using the bacterium Streptococcus gordonii which can express heterologous Ag and can colonize the oral cavity and vagina of mice 268-D and 257-D recognized S. gordonii expressing the V3 domain of MN the vaccine stimulated V3-specific IgG2a in mice. Oggioni et al. [1999]
- 268-D: Review of clade specificity and anti-V3 HIV-1-Abs. Zolla-Pazner *et al.* [1999a]
- 268-D: Peptide reactivity pattern clustered with immunological related MAbs: 1108, 386, 268, 311, 257, 694.8 the amino acids HI tended to be critical for reactivity in this group MAb 453, with an identical core epitope to 268 based on prior experiments (HIGPGR), was not part of this reactivity group, illustrating that context can be critical. Zolla-Pazner *et al.* [1999b]
- 268-D: A T-cell line-adapted (TCLA) derivative of SI primary isolate 168P acquired the ability to be neutralized by anti-V3 MAbs – the primary isolate could use either CCR5 or CXCR4, and was not neutralized when infection was directed via either pathway, however the TCLA derivative uses CXCR4 only and is neutralized. LaCasse et al. [1998]
- 268-D: Poor reactivity against HIV-1 isolates SF162 and SF128A and no neutralization, in contrast to MAbs 391/95-D and 257-D. Stamatatos et al. [1997]
- 268-D: Failed to neutralize HXB2 and chimeric virus with gp120 from primary isolates in an HXB2 background. McKeating et al. [1996]
- 268-D: 268-D is V H4 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996]

- anti-V3, and anti-CD4BS MAbs to monomeric and virionassociated gp120 from HIV-1 isolates with differences in cell tropism was studied - V3 loop epitopes were less accessible to Ab binding on the virion surface than in the gp120 monomer, particularly for macrophage-tropic isolates SF162 and SF128a, relative to T-cell tropic SF2 – sCD4 association with gp120 did not influence the binding of 268-D to virion-associated gp120, although sCD4 binding did alter epitope exposure for other anti-V3 MAbs. Stamatatos & Cheng-Mayer [1995]
- 268-D: Serotyping study using flow-cytometry, if H of HIGPGR was substituted in virus, 268-D did not bind. Zolla-Pazner et al. [1995]
- 268-D: Moderate dissociation rate and homologous neutralization titer. VanCott et al. [1994]
- 268-D: Neutralizes MN binds SF2: YIGPGR specificity: MN, SF2, NY5, RF, CDC4. Gorny et al. [1993]
- 268-D: Mediated deposition of complement component C3 on HIV infected cells, but not in the presence of sCD4. Spear et al.
- 268-D: Reacts with MN, NY5, CDC4, RF and SF2, does not cross-react with WM52 or HXB2. Karwowska et al. [1992b]
- 268-D: Called 268-11-D-IV strain specific weakly neutralizing. D'Souza et al. [1991]

No. 527

**MAb ID** 386-D (386, 386-10D, 386D)

HXB2 Location gp160 (310-315)

Author Location gp120 (MN)

Epitope HIGPGR

Subtype B

Neutralizing L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zol-Med.

las01@mcrcr6.med.nyu) (NYU

Center)

References Gorny et al. 2004; Gorny & Zolla-Pazner

2004; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Fontenot et al. 1995; VanCott et al. 1994; Gorny et al.

1993; Karwowska et al. 1992b

Keywords antibody binding site definition and exposure, binding affinity, isotype switch, review, sub-

type comparisons

- 386-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 386-D: Called 386: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides do not show as much ability to neutralize as V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. 386 was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)

• 268-D: The binding of conformation-dependent anti-V2, • 386-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H – 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding – V3 MAbs tended to have the most crossreactive binding to clade A, B, C, and D isolates, less to E, F, G, and H – 386-D showed intermediate reactivity. Nyambi et al. [2000] (isotype switch, subtype comparisons)

- 386-D: Review of clade specificity and anti-V3 HIV-1-Abs. Zolla-Pazner *et al.* [1999a] (**review**, **subtype comparisons**)
- 386-D: Peptide reactivity pattern clustered with immunological related MAbs: 1108, 386, 268, 311, 257, 694.8 - the amino acids HI tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure)
- 386-D: Slow dissociation rate, potent homologous neutralization. VanCott et al. [1994] (binding affinity)
- 386-D: Neutralizes MN binds SF2: YIGPGR specificity: MN, SF2, NY5, RF, CDC4. Gorny et al. [1993] (antibody binding site definition and exposure)

No. 528

**MAb ID** 5042A

HXB2 Location gp160 (310–315)

**Author Location** gp120 (310–315 BH10)

Epitope QrGPGR

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: B clade BH10

HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Gorny et al. 1991; Langedijk et al. 1991

• 5042A: Generation and fine mapping of murine MAbs. Langedijk et al. [1991]

No. 529

**MAb ID** 5042B

HXB2 Location gp160 (310–315)

**Author Location** gp120 (310–315 BH10)

Epitope QRGPGr

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade BH10

HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Langedijk et al. 1991

• 5042B: Generation and fine mapping of murine MAbs. Langedijk et al. [1991]

**No.** 530

**MAb ID** 418-D (418, 418D)

HXB2 Location gp160 (310-316)

Author Location gp120 (MN)

Epitope HIGPGRA

Subtype B

Neutralizing L

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 V3

Center)

References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Zhang et al. 2002; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Gorny et al. 1993; Karwowska et al. 1992b

**Keywords** antibody binding site definition and exposure, review, subtype comparisons, variant cross-recognition or cross-neutralization

- 418-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 418-D: Called 418: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. 418 was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 418-D: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002]
- 418-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding V3 MAbs tended to have the most cross-reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H 418-D showed intermediate reactivity. Nyambi *et al.* [2000] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 418-D: Review of clade specificity and anti-V3 HIV-1-Abs. Zolla-Pazner *et al.* [1999a] (**review**, **subtype comparisons**)
- 418-D: Called 418 MAb peptide-reactivity pattern clustered with immunological related MAbs: 391.5, 412 and 418, all selected with MN V3 peptide the core amino acids HIGPGR tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure)
- 418-D: Neutralizes MN, does not bind to SF2 or HXB2. Gorny et al. [1993] (variant cross-recognition or cross-neutralization)
- 418-D: MN strain specific, does not cross-react with SF2, NY5, RF, CDC4 WM52 or HXB2. Karwowska *et al.* [1992b] (variant cross-recognition or cross-neutralization)

**No.** 531 **MAb ID** 5021

**HXB2 Location** gp160 (310–316)

Author Location gp120

Epitope QrGPGRa

Neutralizing  $\,L\,$ 

Immunogen vaccine

Vector/Type: peptide Strain: B clade BH10

HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Moore et al. 1993b; Langedijk et al. 1991; Durda et al. 1990; Durda et al. 1988

- 5021: Binding to native gp120 100-300 fold greater than to denatured 314G/W substitution abolishes binding, changes outside the loop have little effect. Moore *et al.* [1993b]
- 5021: Generation and fine mapping of murine MAbs. Langedijk *et al.* [1991]

No. 532

**MAb ID** 5025B

HXB2 Location gp160 (310-316)

**Author Location** gp120 (310–316 BH10)

Epitope QRGPGra

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade BH10

HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Langedijk et al. 1991

• 5025B: Generation and fine mapping of murine MAbs. Langedijk *et al.* [1991]

**No.** 533

**MAb ID** 5042

**HXB2 Location** gp160 (310–316)

Author Location gp120

Epitope QRGPGRA

Neutralizing L

Immunogen vaccine

Vector/Type: peptide

Species (Isotype) mouse

Ab Type gp120 V3

References Moore et al. 1993b; Durda et al. 1990; Durda

et al. 1988

• 5042: Binding to native gp120 100-300 fold greater than to denatured – 314G/W substitution abolishes binding, changes outside the loop have little effect. Moore *et al.* [1993b]

**No.** 534

**MAb ID** 110.3

HXB2 Location gp160 (310-317)

Author Location gp120 (308–328 BRU)

Epitope QRGPGRAF

Neutralizing L

Immunogen vaccine

Vector/Type: HIV infected-cell lysate Strain: B clade BRU HIV component:

HIV-1

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type gp120 V3

References Connelly et al. 1994; Pirofski et al. 1993; • 110.4: 313 P/S substitution in the V3 region disrupts binding. Langedijk et al. 1992; Evans et al. 1989; Thomas et al. 1988

- 110.3: An anti-idiotypic MAb generated against 110.3 both mimics and binds to V3, suggesting that the V3 loop may associated with itself. Connelly et al. [1994]
- 110.3: MAb variable region sequenced heavy chain: V 7138(40), D deletion, J H4 – light chain: V kappa21(47), J kappa2. Pirofski et al. [1993]
- 110.3: Included as a control. Evans et al. [1989]

No. 535 **MAb ID** 110.4

HXB2 Location gp160 (310-317)

Author Location gp120 (308-328 BRU)

Epitope QRGPGRAF

Neutralizing L

Immunogen vaccine

HIV infected-cell lysate *Vector/Type:* Strain: B clade BRU HIV component: HIV-1

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type gp120 V3

Research Contact Genetic Systems Corp, Seattle WA, E. Kinney-Thomas

References Guillerm et al. 1998; Cao et al. 1997b; Valenzuela et al. 1998; McDougal et al. 1996; Connelly et al. 1994; Boudet et al. 1994; Thali et al. 1994; Arendrup et al. 1993; Pirofski et al. 1993; Thali et al. 1993; Langedijk et al. 1992; Thali et al. 1992b; Callahan et al. 1991; Thomas et al. 1988

**Keywords** anti-idiotype, antibody binding site definition and exposure, antibody sequence, variable domain, escape

- 110.4: Used for flow cytometry in a study of the anti-CD4, CDR3 loop MAb called 13B8.2, in a study of HIV-1 induced programmed cell death. Guillerm et al. [1998]
- 110.4: Neutralization of LAI in CEM cells by anti-V3 MAbs 110.4 and N11-20 is through inhibition of viral binding to the cell. Valenzuela et al. [1998]
- 110.4: Virus with the V1-V2 loop deleted was viable and more susceptible to neutralization by CD4i MAb 17b, and anti-V3 MAbs 1121, 9284, and 110.4, but not to and CD4BS MAb F105 or sCD4. Cao et al. [1997b] (antibody binding site definition and exposure)
- 110.4: Neutralizes HIV-1 LAI. McDougal et al. [1996]
- 110.4: An anti-idiotypic MAb generated against 110.3 also blocks binding of 110.4. Connelly et al. [1994] (anti-idiotype)
- 110.4: gp41 mutation that confers resistance to neutralization by anti-CD4 binding site antibodies does not reduce neutralizing efficiency of this V3 region MAb. Thali et al. [1994] (antibody binding site definition and exposure)
- 110.4: Primary isolates from different time points from one individual were not susceptible to neutralization by 110.4. Arendrup et al. [1993]
- 110.4: MAb variable region sequenced heavy chain: V 3660-SB32, D closest to DSP2.3, 2.4 and .6, J H2 – light chain: V kappa21, J kappa2. Pirofski et al. [1993] (antibody sequence, variable domain)

Thali et al. [1992b] (antibody binding site definition and exposure, escape)

• 110.4: Polyanionic polysaccharides were proposed to inhibit viral functions such as binding and syncytia formation through interations mediated through the local high positive charge density in the V3 loop. The binding of this anti-V3 antibody is inhibited by dextransulfate. Callahan et al. [1991]

No. 536

**MAb ID** 110.5

HXB2 Location gp160 (310–317)

Author Location gp120 (308-328 BRU)

Epitope QRGPGRAF

**Neutralizing** L

Immunogen vaccine

Vector/Type: HIV infected-cell lysate Strain: B clade BRU HIV component:

HIV-1

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type gp120 V3

Research Contact E. Kinney-Thomas or Genetic Systems, Seattle WA

> References Parren et al. 1998a; Ugolini et al. 1997; Binley et al. 1997a; Jeffs et al. 1996; McDougal et al. 1996; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; Sattentau et al. 1995; Klasse et al. 1993a; Thali et al. 1993; Moore et al. 1993b; Pirofski et al. 1993; McKeating et al. 1992a; Langedijk et al. 1992; Sattentau & Moore 1991; Cordell et al. 1991; Moore et al. 1990; Thomas et al. 1988; Reitz et al. 1988

- 110.5: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated - authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- 110.5: Viral binding inhibition by 110.5 was correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5) Ugolini et al. [1997]
- 110.5: Deletion of the V1V2 regions did not affect anti-V3 Abs ability to bind when compared to intact rec gp120. Jeffs et al. [1996]
- 110.5: Neutralizes HIV-1 LAI. McDougal et al. [1996]
- 110.5: Reciprocal binding inhibition with other anti-V3 MAbs - enhances binding of some anti-V2 MAbs - binding enhanced by some CD4 binding site MAbs. Moore & Sodroski [1996]
- 110.5: Did not induce dissociation of gp120, as sCD4 did discrepancy with Poignard et al. [1996a], that was suggested to be due to MAb interference with detection, as the gp120-MAb complex was denatured in the Poignard study Moore et al. [1990]. Moore et al. [1990]; Poignard et al. [1996a]
- 110.5: V3 MAbs 9284, BAT123, 110.5, and 110.I could each significantly increase gp120 dissociation from virus, mimicking sCD4, and expose the gp41 epitope for MAb 50-69, in contrast to anti-V2 MAbs. Poignard et al. [1996a]
- 110.5: Pretreatment of HX10-infected H9 cells with sCD4 decreases signal from 110.5 at 37 degrees due to dissociation of gp120-gp41. Sattentau et al. [1995]

- 110.5: Binds with high affinity to monomer and oligomer, rapid association and potent neutralization of lab strains - neutralizes cell-free Hx10. Sattentau & Moore [1995]
- 110.5: The gp41 mutation 582(Ala to Thr) results in conformational changes in gp120 that confer neutralization resistance to conformationally sensitive neutralizing MAbs - neutralization efficiency of 110.5 is not affected. Klasse et al. [1993a]; Reitz et al. [1988]
- 110.5: Thrombin cleavage of V3 loop between R-315 and A-316 abrogates binding – can inhibit C4 region antibody which has conformational requirements (G3-299) - binding to native gp120 100-300 fold greater than to denatured. Moore et al. [1993b]
- 110.5: Variable region sequenced heavy chain: V 3660-SB32, D closest to DSP2.3, 2.4 and .6, J H2 – light chain: V kappa21, J kappa2. Pirofski et al. [1993]
- 110.5: Binding insensitive to gp120 reduction. Cordell et al. [1991]
- 110.5: Two fold increase in binding to gp120 in the presence of bound sCD4. Sattentau & Moore [1991]

**No.** 537

**MAb ID** 58.2

HXB2 Location gp160 (310-317)

Author Location gp120 (MN)

Epitope HIGPGRAF

Neutralizing L

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN

HIV component: V3

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type gp120 V3

Research Contact Repligen Corp.

References Binley et al. 2004; York et al. 2001; Stanfield et al. 1999; Seligman et al. 1996; Moore et al. 1994b; Potts et al. 1993; White-Scharf et al. 1993

Keywords subtype comparisons, recognition or cross-neutralization

- 58.2: 93 viruses from different clades were tested for their neutralization cross-reactivity using a panel of HIV antibodies. 58.2 could only neutralize B subtype viruses, and seemed to have a minimal epitope of (H/T)IGPGR(A/T)(F/L). Binley et al. [2004] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 58.2: 58.2's epitope was noted to be IGPGRAF Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived Research Contact Susan from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding. York et al. [2001]
- 58.2: The crystal structure of Fab 58.2 bound to V3 loop peptides was obtained - conformational changes in the tip of the V3 loop (GPGR) were observed when different MAbs were bound – 58.2's epitope was defined as KRKRIHIGPGRAFY. Stanfield et al. [1999]

- 58.2: Competition ELISAs with serial deletions produced longer estimates of epitope length, RIHIGPGRAFY, than Alanine substitution, suggesting significance of non-contact residues. Seligman et al. [1996]
- 58.2: Modest cross-reactivity among B clade gp120s, little outside B clade – core epitope as I-IHIG. Moore et al. [1994b]
- 58.2: Did not synergistically neutralize MN in combination with MAb F105 – there was synergistic neutralization when combined with sCD4. Potts et al. [1993]
- 58.2: Epitope defined by peptide reactivity and changes in affinity with amino acid substitutions – 4/7 primarily isolates were neutralized. White-Scharf et al. [1993]

No. 538

MAb ID polyclonal

HXB2 Location gp160 (310-318)

Author Location gp120

Epitope QRGPGRAFV?

Neutralizing L

Immunogen vaccine

Vector/Type: peptide keyhole limpet hemocyanin (KLH) conjugate, peptide Brucella abortus (Ba) conjugate, peptide lipopolysaccharide (LPS) conjugate Strain: B clade MN HIV component: V3

Species (Isotype) mouse (IgA, IgG1, IgG2a)

References Golding et al. 2002a

• Internasal (i.n.) immunization with V3-Ba induced mucosal anti-V3 NAbs and IFN-gamma secreting T cells - V3-Ba, V3-KLH and V3-LPS could each induce serum and mucosal IgA and IgG in BALB/c mice - i.n. plus i.p. immunizations gave higher titers than i.n. alone - the response to V3-KLH was mainly restricted to IgG1, and to V3-Ba, IgG2a - class II KO mice (CD4+-deficient) did not respond to V3-KLH, but did respond to V3-Ba, suggesting that V3-Ba may be effective in eliciting Ab responses in HIV-1 infected individuals that have impaired CD4+ T cell function. Golding et al. [2002a]

No. 539

**MAb ID** 537-D (537)

HXB2 Location gp160 (311–315)

**Author Location** gp120 (MN)

Epitope IGPGR

Subtype B

**Neutralizing** L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

Zolla-Pazner (Zollas01@mcrcr6.med.nyu) Med.

Center)

References Gorny et al. 2004; Nyambi et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Fontenot et al. 1995; VanCott et al. 1994; Gorny et al. 1993; Gorny et al. 1992; Kar-

wowska et al. 1992b

**Keywords** antibody binding site definition and exposure

- 537-D: Called 537: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 537-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H - 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding - V3 MAbs tended to have the most crossreactive binding to clade A, B, C, and D isolates, less to E, F, G, and H – 537-D showed weak reactivity. Nyambi et al. [2000]
- 537-D: Review of clade specificity and anti-V3 HIV-1-Abs. Zolla-Pazner et al. [1999a]
- 537-D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 1334, 419, 504, 447, 453 and 537 - the core amino acids GP tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b]
- 537-D: Moderate homologous neutralization, relatively rapid Research Contact Paul Durda, Du Pont de Nemours and Co dissociation constant. VanCott et al. [1994]
- 537-D: MN type specific neutralization observed binds SF2, also IGPGR. Gorny et al. [1992, 1993]
- 537-D: Reacts with MN, NY5, CDC4, RF, WM52 and SF2, but does not cross-react with HXB2. Karwowska et al. [1992b]

**No.** 540

**MAb ID** 5020

HXB2 Location gp160 (311–316)

**Author Location** gp120 (311–316 BH10)

Epitope RGPGRA

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade BH10

HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Langedijk et al. 1991

• 5020: Generation and fine mapping of murine MAbs. Langedijk et al. [1991]

No. 541

MAb ID RC25

HXB2 Location gp160 (311-316)

Author Location gp120 (JRFL)

Epitope IGPGRA

Subtype B

Neutralizing L

**Immunogen** 

Species (Isotype) humanized mouse

Ab Type gp120 V3

References Kaizu et al. 2003; Kimura et al. 2002

Keywords co-receptor, HAART, ART

• RC25: MD14 is a R5X4 SHIV with a B clade Env; the V3 loop of an E-clade Env was inserted into MD14 to create SHIV-TH09V3, an R5 virus. SHIV-TH09V3 could infect both cynomolgous and pig-tailed macaques, and the R5 co-receptor usage was maintained after passage through macaques. The MAb RC25 recognized B clade V3 loops, and reacted with

SHIV-MD14. Rabbit anti-sera raised against a NSI Clade E consensus preferentially recognized SHIV-TH09V3. Kaizu et al. [2003] (co-receptor)

RC25: RC25 is a humanized MAb that recognizes the epitope IGPGRA - it has strong neutralizing activity against JRFL (R5 virus) and weak against NL4-3 (X4 virus) and is used as a control in a study of NAb activity in patients undergoing HAART. Kimura et al. [2002] (HAART, ART)

No. 542

MAb ID 5023A (5023, NEA-9205, NEA 9205)

**HXB2 Location** gp160 (311–317) **Author Location** gp120 (311–317 BH10)

Epitope RgPGRAF

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: B clade BH10 HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Schonning et al. 1998; Rovinski et al. 1995;

Back et al. 1993; D'Souza et al. 1991; Langedijk et al. 1991

- 5023A: Called NEA-9205 The N306 glycan of the V3 loop makes the tip of the V3 loop inaccessible to this MAb in oligomeric Env, loss of this glycan enhances neutralization sensitivity. Schonning et al. [1998]
- 5023A: Called 5023 in this paper Used to precipitate gp160 in immunoblots in a study examining the feasibility of using unprocessed gp160 glycoprotein as an immunogen. Rovinski
- 5023A: Called 5023 Langedijk also has an MAb called 5023B - gp41 amino acid substitutions 668 (N/S) and 675 (I/M) in gp41 interfere with 5023s neutralization potency, region 662-675 is ELDKWANLWNWFNI. Back et al. [1993]
- 5023A: Called 5023 Langedijk also has an MAb called 5023B - strong cross-reactive neutralizing MAb. D'Souza et al. [1991]
- 5023A: Generation and Fine mapping of murine MAbs. Langedijk et al. [1991]

No. 543

**MAb ID** 110.6

**HXB2 Location** gp160 (311–318)

Author Location gp120 (BRU)

**Epitope** RGPGRAFV

**Neutralizing** L (weak)

Immunogen vaccine

HIV infected-cell lysate Vector/Type: Strain: B clade BRU HIV component:

HIV-1

**Species (Isotype)** mouse (IgG1 $\lambda$ )

Ab Type gp120 V3

References Langedijk et al. 1992; Pirofski et al. 1993;

Thomas et al. 1988

• 110.6: Variable region sequenced - heavy chain: V J558-146b.1alpha, D closest to DSP16.2, J H3 – light chain: V lambda1, J lambda1. Pirofski et al. [1993]

No. 544

MAb ID polyclonal HXB2 Location gp160 (311–318)

Author Location gp120 (MN)

Epitope IGPGRAFY

Neutralizing L

Immunogen vaccine

Vector/Type: B. abortus complex Strain: B clade MN, B clade SF2 HIV component:

gp120

Species (Isotype) mouse (IgG2a)

Ab Type gp120 V3

References Golding et al. 1995

• Ab is evoked even in mice depleted of CD4+ cells.

No. 545

MAb ID 10/36e

**HXB2 Location** gp160 (311–321)

Author Location gp120 (311–321 HXB10)

Epitope RGPGRAFVTIG Neutralizing L (HXB10) Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2a)

Ab Type gp120 V3

**References** Peet *et al.* 1998; McKeating *et al.* 1993b; McKeating *et al.* 1992a

- 10/36e: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind, but anti-V3 MAb 10/36e binding was dramatically diminished by V3 serine substitutions mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet *et al.* [1998]
- 10/36e: Binding to virion gp120 enhanced by sCD4. McKeating *et al.* [1992a]

No. 546

**MAb ID** 10/54 (10/54ow/6i/6i)

HXB2 Location gp160 (311–321)

Author Location gp120 (311–321 HXB10)

Epitope RGPGRAFVTIG Neutralizing L (HXB10) Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG1)

Ab Type gp120 V3

**References** Peet *et al.* 1998; McKeating *et al.* 1993b; McKeating *et al.* 1993a; McKeating *et al.* 

1992a

• 10/54: Called 10/54ow/6i/6i: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic – these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind, but anti-V3 MAb 10/54 binding was dramatically diminished by V3 serine substitutions – mice injected with serine substituted gp120 had a reduced response relative to wildtype,

and no enhanced immunogenicity of conserved regions. Peet et al. [1998]

- 10/54: Studied in the context of a neutralization escape mutant. McKeating et al. [1993a]
- 10/54: Binding to virion gp120 enhanced by sCD4. McKeating et al. [1992a]

No. 547

**MAb ID** 11/85b (11/85b/14I/14I)

HXB2 Location gp160 (311–321)

Author Location gp120 (311-321 HXB10)

**Epitope** RGPGRAFVTIG **Neutralizing** L (HXB2)

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2b)

**Ab Type** gp120 V3

**References** McKeating *et al.* 1993b; McKeating *et al.* 

1992a

11/85b: Binding to virion gp120 enhanced by sCD4. McKeating et al. [1992a]

**No.** 548

MAb ID polyclonal

HXB2 Location gp160 (311-322)

**Author Location** gp120 (MN)

Epitope IGPGRAFYTTKN

**Neutralizing** L (MN ALA-1)

Immunogen vaccine

Vector/Type: human rhinovirus 14 Strain: B

clade MN HIV component: V3

**Species (Isotype)** guinea pig **Ab Type** gp120 V3

References Smith et al. 1998

• The tip of the MN V3 loop (IGPGRAFYTTKN) was inserted into cold-causing human rhinovirus 14 (HRV14) – chimeras were immunoselected, and chimeric viruses were neutralized by anti-V3 loop antibodies – chimeric viruses elicited potent NAbs against ALA-1 and MN. Smith *et al.* [1998]

No. 549

**MAb ID**  $0.5\beta$  (0.5 beta, 0.5beta)

HXB2 Location gp160 (311-324)

Author Location gp120 (316–330 HXB2)

Epitope RGPGRAFVTIGKIG

Subtype B

Neutralizing L (IIIB)

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: Env

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type gp120 V3

Research Contact Shuzo Matsushita or Toshio Hattori of Ku-

mamoto University

References Harada et al. 2004; Kawai et al. 2003; Zvi et al. 2000: Tugarinov et al. 2000: Jagodzinski & Trzeciak 2000; Fortin et al. 2000; Tugarinov et al. 1999; Faiman & Horovitz 1997; Wyatt et al. 1997; Zvi et al. 1997; Huang et al. 1997; Faiman et al. 1996; Jeffs et al. 1996; McDougal et al. 1996; Warrier et al. 1996; Jagodzinski et al. 1996; Zvi et al. 1995a; Zvi et al. 1995b; Broder et al. 1994; Boudet et al. 1994; Okada et al. 1994; Thali et al. 1994; Cook et al. 1994; Watkins et al. 1993; Klasse et al. 1993a; Moore et al. 1993b; di Marzo Veronese et al. 1993; Sperlagh et al. 1993; McKeating et al. 1992a; Maeda et al. 1992; Emini et al. 1992; Matsushita et al. 1992; D'Souza et al. 1991; Nara et al. 1990; Reitz et al. 1988; Skinner et al. 1988a; Skinner et al. 1988b; Matsushita et al. 1988

**Keywords** anti-idiotype, antibody binding site definition and exposure, antibody generation, antibody interactions, brain/CSF, co-receptor, complement, escape, structure, variant cross-recognition or cross-neutralization

- 0.5beta: UK Medical Research Council AIDS reagent: ARP3025.
- 0.5beta: NIH AIDS Research and Reference Reagent Program: 1591.
- 0.5beta: Studies on the temperature dependence of infectious virus (increased temperatures up to 37 degress increases infectivity) showed that X4 pseudoviruses that were infectious at room temperature were also more resistant to anti-V3 0.5beta and anti-CXCR4 blocking peptide T140. This implies that virus more heavily populated with functional envelopes are more infectious. Harada *et al.* [2004] (**co-receptor**)
- 0.5beta: 0.5beta was used as a control for gp120 expression relative to Nef expression soon after infection of cultures. The goal of this study was to elicit anti-Nef IgM human Abs in trans-chromosome mice that carry human chromosomes responsible for Ig production. These mice were immunized with recombinant Nef protein. Human heavy chain, mouse light chain anti-Nef IgM were obtained. The hope was that these human IgM anti-Nef chimeric Abs would activate complement mediated cytolysis; C3 deposition was induced but not at high enough levels to induce lysis. Kawai *et al.* [2003] (complement)
- 0.5beta: Host encoded intercellular adhesion molecule (ICAM-1) is incorporated by the HIV-1 virion and enhances viral infectivity ICAM-1 does not modify virus sensitivity to antibodies 0.5beta or 4.8D or sCD4, but neutralizing ability of F105 was diminished in ICAM bearing virions in the presence of lymphocyte function-association antigen-1 (LFA-1) Ab. Fortin *et al.* [2000] (antibody interactions)
- 0.5beta: MAbs 0.5beta and G3-42 were used to study synthesis of oligomeric and monomeric forms of Env inhibition of glycosylation by tunicamycin results in reduction of oligomeric gp120 at the cell surface and of monomer in the cytoplasm neither MAb recognized non-glycosylated Env precursor. Jagodzinski & Trzeciak [2000]

0.5beta: 14/18 residues of peptide P1053, RKSIRIQRGP-GRAFVTIG, were shown to be involved in the Ab recognition site using NMR – QRGPGR forms a beta-hairpin turn at the center of the binding pocket. Tugarinov *et al.* [2000] (antibody binding site definition and exposure)

- 0.5beta: NMR and mutation cycles were employed to generate a model of the peptide-antibody complex, showing aa residues that interact or do not contribute to the binding of MAb 0.5beta Fv with the peptide F96(L) of 0.5beta binds to Pro13, H52(H) interacts with Ile7, Ile9, Gln10, and D56(H) interacts with Arg11 of the V3 loop peptide RGPG retains hairpin conformation binds in the center of a groove. Zvi *et al.* [2000] (**structure**)
- 0.5beta: NMR structure reveals that Ab bound IIIB-V3 peptide adopts an unexpected type VI cis proline beta-turn. Tugarinov et al. [1999] (structure)
- 0.5beta: The Fv fragment was purified and the temperature dependence and effect of mutations was studied. Faiman & Horovitz [1997]
- 0.5beta: Relative to the native peptide, an O-linked alphagalactosamine modified V3 peptide enhanced binding to 0.5 beta, while an N-linked beta-glucosamine modified peptide showed reduced binding. Huang et al. [1997] (antibody binding site definition and exposure)
- 0.5beta: Binds both gp120 and soluble gp120+gp41 complex efficiently, suggesting its gp120 epitope is not blocked by gp41 binding. Wyatt *et al.* [1997] (antibody binding site definition and exposure)
- 0.5beta: The structure of a 17 amino acid V3 peptide bound to the Fab was studied using NMR. Zvi et al. [1997] (structure)
- 0.5beta: For Fv fragment of 0.5beta, the combined variable regions of the heavy and light chain residues, were purified. Binding of the V3 peptide epitope TRKSIRIQRGPGRAFVTIGK was studied through mutagenesis of arginines and the free energy of binding in various salt concentrations. R4A, R8A, and R11A all reduce the free energy; R8 is embedded in the peptide-Fv fragment, while R11 is more solvent exposed. Faiman *et al.* [1996] (antibody binding site definition and exposure)
- 0.5beta: The sulfated polysaccharide curdlan sulfate (CRDS) binds to the Envelope of T-tropic viruses and neutralizes virus CRDS inhibits 0.5beta binding 0.5beta epitope described as GPGRAFVTIG. Jagodzinski *et al.* [1996]
- 0.5beta: Deletion of the V1V2 regions did not affect anti-V3
   Abs ability to bind when compared to intact rec gp120. Jeffs
   et al. [1996] (antibody binding site definition and exposure)
- 0.5beta: Synergistic neutralization of HIV-1 when combined with anti-V2 MAb C108G. Warrier *et al.* [1996] (antibody interactions)
- 0.5beta: The interactions of the peptide RKSIRIQRGP-GRAFVT 0.5beta were studied by NMR, and hydrophobic interactions between the two Is and the V form the base of a 12 amino acid loop with GPGR at the apex. Zvi *et al.* [1995b] (antibody binding site definition and exposure)
- 0.5beta: NMR of 0.5beta bound NNTRKSIRIQRGP-GRAFVTIGKIG suggests that the bound amino acids are in the region SIRIQRGPGRAFVT. Zvi *et al.* [1995a] (antibody binding site definition and exposure)

- 0.5beta: Type-specific neutralization of IIIB does not neutralize SF2. Broder et al. [1994] (variant cross-recognition or cross-neutralization)
- 0.5beta: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon – this MAb can inhibit gp120 binding to GalCer in vitro. Cook et al. [1994] (brain/CSF)
- 0.5beta: Binding domain aa 310-319: RGPGRAFVTIGKIG - mutations in the V3 loop from basic residues can destroy virus infectivity and syncytium formation: R306T, R309T and R313G can also reduce binding of V3 MAbs with two different binding sites: 9284 and 0.5beta. Okada et al. [1994] (antibody binding site definition and exposure)
- 0.5beta: gp41 mutation that confers resistance to neutralization by anti-CD4 binding site antibodies does not reduce neutralizing efficiency of this V3 region MAb. Thali et al. [1994]
- 0.5beta: The gp41 mutation 582(Ala to Thr) results in conformational changes in gp120 that confer neutralization resistance to some antiserum and conformationally sensitive neutralizing MAbs – neutralization efficiency of 0.5beta is not affected. Klasse et al. [1993a]; Reitz et al. [1988] (antibody binding site definition and exposure)
- 0.5beta: Binding to native gp120 100-300 fold greater than to denatured. Moore et al. [1993b] (antibody binding site definition and exposure)
- 0.5beta: Monoclonal anti-idiotype antibodies that mimic the 0.5beta epitope were generated. Sperlagh et al. [1993] (antiidiotype)
- 0.5beta: A neutralization escape mutant (HXB2 A281V) was selected by growth of HXB2 in the presence of broadly neutralizing sera - of the MAbs tested, 0.5beta neutralization was the most profoundly affected by this mutation. Watkins et al. [1993] (escape)
- 0.5beta: Neutralization of virus carrying an A to T substitution Research Contact M. Terada, Jason Grabely (contrast with MAb M77) di Marzo Veronese et al. [1993]
- 0.5beta: sCD4 causes loss of IIIB type-specificity, allowing binding and neutralization of MN, in contrast to MAb mu5.5. Maeda et al. [1992]
- 0.5beta: Chimeric mouse-human MAb Cbeta1 was constructed by combining the human Cgamma1 and Ckappa constant regions with the 0.5beta murine MAb – ADCC and neutralizing activity. Matsushita et al. [1992] (complement)
- 0.5beta: Potent neutralizing activity. D'Souza et al. [1991]
- 0.5beta: Emergence of virus resistant to MAb 0.5beta and autologous sera neutralization in IIIB infected chimps. Nara et al. [1990] (escape)
- 0.5beta: Type-specific neutralization of IIIB does not neutralize MN or RF. Matsushita et al. [1988]; Skinner et al. [1988b] (antibody generation)

**No.** 550

**MAb ID** C $\beta$ 1, 0.5 $\beta$ 

HXB2 Location gp160 (311–324)

**Author Location** gp120 (316–330 HXB2)

Epitope RGPGRAFVTIGKIG

Subtype B

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: Env

**Species (Isotype)** humanized mouse (IgG1)

Ab Type gp120 V3

References Ferrantelli & Ruprecht 2002; Kimura et al. 2002; Matsushita et al. 1992; Emini et al.

- Cbeta1: Review of passive immunoprophylaxis with human NAbs that also includes this chimeric mouse-human MAb, noting it protected 2/2 Chimpanzees from HIV-1 IIIB infection in the Emini et al. study published in 1992. Ferrantelli & Ruprecht [2002]
- Cbeta1: Defines epitope as IQRGPGRA strong neutralizing activity against NL4-3 (X4 virus) and none against JRFL (R5 virus) – used as a control in a study of NAb activity in patients undergoing HAART. Kimura et al. [2002]
- Cbeta1: passive transfer to chimpanzees confers protection against challenge with homologous cell-free virus - mouse 0.5beta human IgG1 chimera. Emini et al. [1992]
- Cbeta1: Chimeric mouse-human MAb Cbeta1 was constructed by combining the human Cgamma1 and Ckappa constant regions with the 0.5beta murine MAb – ADCC and neutralizing activity. Matsushita et al. [1992]

No. 551

MAb ID NM-01 (hNM01, hNM-01)

HXB2 Location gp160 (312–315)

Author Location gp120 (MN)

Epitope GPGR

**Neutralizing** L

Immunogen vaccine

Vector/Type: human rhinovirus 14 Strain: B clade MN HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Zwick et al. 2003; Nakamura et al. 2000; Smith et al. 1998; Yoshida et al. 1997; Ohno

et al. 1991

Keywords antibody interactions, complement, immunotherapy

- NM-01: Called hNM01. scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. The humanized version of this MAb was one of the V3 MAbs used. Zwick et al. [2003] (antibody interactions)
- NM-01: Called hNM01. The CDR region of the murine MAb NM-01 was put into a human IgG frame. The epitope recognition was preserved, but the neutralizing potency of the humanized form was enhanced. It could activate complement. Nakamura et al. [2000] (complement, immunotherapy)

NM-01: The tip of the MN V3 loop was inserted into cold-causing human rhinovirus 14 (HRV14) – chimeras were immunoselected, and chimeric viruses were neutralized by anti-V3 loop antibodies, and NM-01 was among the Abs used – chimeric viruses elicited potent NAbs in guinea pigs against ALA-1 and MN. Smith et al. [1998]

NM-01: Resistance mutation selected by propagation of molecular cloned isolate in the presence of NM-01. Yoshida *et al.* [1997]

No. 552

**MAb ID** 1026

HXB2 Location gp160 (312-317)

Author Location gp120 (MN)

Epitope GPGRAF

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp120

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Bou-Habib et al. 1994; Nakamura et al. 1993

- 1026: Greater affinity for T cell-tropic strain T-CSF, derived from JR-CSF, than to the primary isolate JR-CSF. Bou-Habib *et al.* [1994]
- 1026: Bound diverse strains, neutralizing activity against MN, close to GPGRAF. Nakamura *et al.* [1993]

No. 553

**MAb ID** 1034

**HXB2 Location** gp160 (312–317)

Author Location gp120 (MN)

Epitope GPGRAF

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

HIV component: gp120

 $\boldsymbol{Species}\;(\boldsymbol{Isotype})\;\;\text{mouse}\;(\boldsymbol{IgG})$ 

Ab Type gp120 V3

References Berman et al. 1997; Bou-Habib et al. 1994

- 1034: Binds to 5/7 isolates from breakthrough cases from a MN gp120 vaccine trial. Berman *et al.* [1997]
- 1034: Greater affinity for T cell tropic T-CSF, derived from JR-CSF, than to the primary isolate JR-CSF, close to GPGRAF. Bou-Habib *et al.* [1994]

No. 554

**MAb ID** 59.1 (R/V3-59.1)

**HXB2 Location** gp160 (312–317)

Author Location gp120 (308-313 MN)

Epitope GPGRAF

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN

HIV component: V3

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

Research Contact Mary White-Scharf and A. Profy, Repligen

Corporation

**References** York *et al.* 2001; Stanfield *et al.* 1999; Smith *et al.* 1998; Ghiara *et al.* 1997; Seligman *et al.* 1996; D'Souza *et al.* 1994; Bou-Habib *et al.* 1994; Ghiara *et al.* 1993; Potts *et al.* 1993; White-Scharf *et al.* 1993; D'Souza *et al.* 1991

- 59.1: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding. York et al. [2001]
- 59.1: The crystal structure of V3 loop peptides bound to Fabs was obtained conformational changes in the tip of the V3 loop (GPGR) were observed when different MAbs were bound. Stanfield *et al.* [1999]
- 59.1: The tip of the MN V3 loop was inserted into cold causing human rhinovirus 14 (HRV14) chimeras were immunoselected, and chimeric viruses were neutralized by anti-V3 loop antibodies, and 59.1 was among the Abs used chimeric viruses elicited potent NAbs in guinea pigs against ALA-1 and MN. Smith *et al.* [1998]
- 59.1: A conformationally restricted analog of the tip of the V3 loop was constructed and bound with Fab 59.1 crystal structure shows interactions between 59.1 and an MN peptide and 59.1 and the modified peptide are similar, but NMR studies reveal that the modified peptide is more ordered in solution, retaining the Fab bound form. Ghiara *et al.* [1997]
- 59.1: Competition ELISAs with serial deletions produced longer estimate of epitope length than x-ray crystallography or Alanine substitution, RIHIGPGRAFYTT, suggesting significance of non-contact residues. Seligman et al. [1996]
- 59.1: Greater affinity for T-cell tropic strain T-CSF than the primary isolate JR-CSF, from which T-CSF was derived. Bou-Habib *et al.* [1994]
- 59.1: Multi-lab study for antibody characterization and assay comparison neutralizes MN and IIIB. D'Souza *et al.* [1994]
- 59.1: Crystal structure of a 24 amino acid peptide from the V3 loop bound to 59.1 Fab fragment – contact residues IGPGRAF. Ghiara et al. [1993]
- 59.1: Synergistic neutralization of MN when combined with sCD4 or the CD4BS MAb F105. Potts *et al.* [1993]
- 59.1: Epitope defined by peptide reactivity and binding affinity with amino acid substitutions – GPGRAF. White-Scharf *et al.* [1993]
- 59.1: Called R/V3-59.1 potent neutralizing MAb. D'Souza et al. [1991]

No. 555

MAb ID polyclonal

**HXB2 Location** gp160 (312–317)

Author Location gp120 (316–321)

Epitope GPGRAF

Neutralizing

Immunogen vaccine

Vector/Type: protein, polyepitope HIV com-

ponent: gp160 Adjuvant: BSA

Species (Isotype) rabbit

**Ab Type** gp120 V3

References Lu et al. 2000b: Lu et al. 2000c

observed upon vaccination with multiple-epitope vaccine CG-GPGRAFY-G-ELDKWA-G-RILAVERYLKD conjugated to BSA, a weak response to GPGRAFY - immunization with CG-(ELDKWA-GPGRAFY)\_2-K was also tried, yielding a strong Ab response to ELDKWA, weak to GPGRAF – gp160 vaccination yielded strong Ab response but not to any of the peptides studied here. Lu et al. [2000c,b]

**No.** 556

**MAb ID** 10E3

HXB2 Location gp160 (312-318) Author Location gp120 (317–323 IIIB)

**Epitope** GPGRAFY

Neutralizing

Immunogen vaccine

Vector/Type: peptide keyhole limpet hemo-

IIIB HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

References Li et al. 2002; Tian et al. 2001

Keywords vaccine antigen design

- 10E3: A polyepitope vaccine was designed based on a recombinant GST fusion protein containing three repeats of the 2F5 core epitope ELDKWA combined with the V3 region peptide GPGRAFY. Abs raised in mice could recognize the peptides, sgp41, and CHO-WT cells that expressed HIV-1 Env on their surface. Li et al. [2002] (vaccine antigen design)
- 10E3: Peptides GPGRAFY and ELDKWAG were conjugated to KLH and used to raise mouse monoclonal Ab - MAb hybridomas were generated with defined specificity – 10E3 binds to the peptide GPGRAFY and to rgp160. Tian et al. [2001]

No. 557

MAb ID polyclonal

HXB2 Location gp160 (312-318)

Author Location gp120 (317-323)

Epitope GPGRAFY

**Neutralizing** 

Immunogen vaccine

Adjuvant: BSA

Species (Isotype) rabbit, mouse

Ab Type gp120 V3

References Yu et al. 2000

• High levels of epitope-specific Abs were induced by the peptide-BSA conjugates C-(GPGRAF)\_4-BSA or C-(TRPNNNTRKSIRIQRGPGRAFYTIG KI)-BSA but not by rgp160 vaccine. Yu et al. [2000]

No. 558

**MAb ID** N11-20 (110-H)

HXB2 Location gp160 (312-320)

Author Location gp120 (317–325)

Epitope GPGRAFVTI

**Neutralizing** L (LAI)

**Immunogen** 

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type gp120 V3

• High titer response to ELDKWA and RILAVERYLKD was Research Contact J. C. Mazie, Hybridolab, Institut Pasteur

References Valenzuela et al. 1998

• N11-20: Neutralization of LAI in CEM cells by anti-V3 MAbs 110.4 and N11-20 is through inhibition of virus binding to the cell. Valenzuela et al. [1998]

**No.** 559

MAb ID 5025A (5025)

HXB2 Location gp160 (313–317)

**Author Location** gp120 (313–317 BH10)

Epitope pgRAF

Neutralizing L

Immunogen vaccine

Vector/Type: peptide Strain: B clade BH10

HIV component: V3

Species (Isotype) mouse (IgG)

Ab Type gp120 V3

cyanin (KLH) conjugate Strain: B clade Research Contact Paul Durda, Du Pont de Nemours and Co

References D'Souza et al. 1991; Langedijk et al. 1991

- 5025: Called 5025 strain specific weakly neutralizing. D'Souza et al. [1991]
- 5025A: Generation and fine mapping of murine MAbs. Langedijk et al. [1991]

**No.** 560

MAb ID N70-1.9b

**HXB2 Location** gp160 (313–318)

**Author Location** gp120 (316–322)

Epitope PGRAFY

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 V3

References Gorny & Zolla-Pazner 2004; Scott et al. 1990;

Robinson et al. 1990a

Keywords ADCC, review, variant cross-recognition or

cross-neutralization

- N70-1.9b: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- Vector/Type: peptide HIV component: V3 N70-1.9b: Type specificity. Robinson et al. [1990a] (variant cross-recognition or cross-neutralization)
  - N70-1.9b: Type specific neutralization, ADCC directed against MN infected cells. Scott et al. [1990] (ADCC, variant crossrecognition or cross-neutralization)

No. 561

**MAb ID** 902

HXB2 Location gp160 (313-324)

Author Location gp120 (IIIB)

Epitope PGRAFVTIGKIG

Neutralizing L

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: gp160

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type gp120 V3

Laboratory, Montana

References Ling et al. 2004; Sakaida et al. 1997; Earl et al. 1994; Broder et al. 1994; Laman et al. 1993; Chesebro & Wehrly 1988

**Keywords** antibody binding site definition and exposure • 902: NIH AIDS Research and Reference Reagent Program:

- 902: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of the V3 MAbs 694-98D and 447-52D, that both bind near the tip of the loop, was decreased by both thrombin and trypsin, 932 binding was only decreased by trypsin. Ling et al. [2004] (antibody binding site definition and exposure)
- 902: V3-BH10 peptide with loop-structure inhibits IL-2 induced T-cell proliferation, thought to be due to altering intracellular signaling, and MAb 908 can block the peptide inhibition. Sakaida *et al.* [1997]
- 902: Epitope may be partially masked or altered in the oligomeric molecule. Broder et al. [1994]
- 902: Used as a control in a study of the influence of oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]
- 902: Strain specific neutralization of HIV. Chesebro & Wehrly [1988]

No. 562

MAb ID 694/98-D (694/98, 694.8, 694/98D)

HXB2 Location gp160 (314-317)

Author Location gp120 (IIIB)

**Epitope** GRAF

Subtype B

Neutralizing L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

Research Contact Dr. Zolla-Pazner, Veterans Affairs Center, NY,

NY. zollas01@endeavor.med.nyu.edu

References Ling et al. 2004; Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Zwick et al. 2003; Zhang et al. 2002; He et al. 2002; Edwards et al. 2002; Park et al. 2000; Nyambi et al. 2000; Altmeyer et al. 1999; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Schonning et al. 1998; Nyambi et al. 1998; Andrus et al. 1998; Li et al. 1998; Smith et al. 1998; Zolla-Pazner et al. 1997; Li et al. 1997; Forthal et al. 1995; Zolla-Pazner et al. 1995; VanCott et al. 1995; Cook et al. 1994; VanCott et al. 1994; Laal et al. 1994; Gorny et al. 1994; Spear et al. 1993; Cavacini et al. 1993a; Gorny et al. 1993; Gorny et al. 1992; Gorny et al. 1991; Skinner et al. 1988b

**Keywords** antibody binding site definition and exposure, antibody interactions, review, variant crossrecognition or cross-neutralization

Research Contact Bruce Chesebro, Rocky Mountain National • 694/98D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)

- 694/98-D: Called 694/98. V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using IIIB gp120. Gorny et al. [2004] (antibody binding site definition and exposure)
- 694-98D: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of the V3 MAbs 694-98D and 447-52D, that both bind near the tip of the loop, was decreased by both thrombin and trypsin. Ling et al. [2004] (antibody binding site definition and exposure)
- 694/98D: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the V3 MAbs used. Zwick et al. [2003] (antibody interactions)
- 694/98-D: Called 694/98D Truncation of the gp41 cytoplasmic domain of X4, R5, and X4R5 viruses forces a conformation that more closely resembles the CD4 bound state of the external Envelope, enhancing binding of CD4i MAbs 17b and 48d and of CD4BS MAbs F105, b12, and in most cases of glycosylation site dependent MAb 2G12 and the anti-gp41 MAb 246D - in contrast, binding of the anti-V2 MAb 697D and the anti-V3 MAb 694/98D were not affected - viruses bearing the truncation were more sensitive to neutralization by MAbs 48d, b12, and 2G12 - the anti-C5 MAb 1331A was used to track levels of cell surface expression of the mutated proteins. Edwards et al. [2002]
- 694/98-D: Called 694 Transgenic mice carrying human genes allowing production of fully human MAbs were used to rapidly create a panel of anti-HIV gp120 MAb producing hybridomas by immunization with HIV SF162 gp120 - the previously described human MAbs 5145A(CD4BS), 4117C (plus others, V3) and 697D (and SC258, V2) were used as controls. He et al.
- 694/98-D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H – 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding - V3 MAbs tended to have the most crossreactive binding to clade A, B, C, and D isolates, less to E, F,

G, and H – 694/98-D showed intermediate reactivity. Nyambi *et al.* [2000]

- 694/98-D: Called 694/98D six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park et al. [2000]
- 694/98-D: A Semliki Forest virus (SFV) expression system carrying BX08 env was used to study the conformation of gp120 intracytoplasmic gp120 was recognized by the anti-V3 MAbs K24 and F5.5, while gp120 at the plasma membrane was detected only by conformation dependent MAbs 2G12, 670-D and 694/98D and not linear V3 MAbs expression in rat brain also showed that surface-expressed Env was recognized only by the conformation-dependent antibodies and not by anti-V3 antibodies. Altmeyer *et al.* [1999]
- 694/98-D: Review of clade specificity and anti-V3 HIV-1-Abs. Zolla-Pazner et al. [1999a]
- 694/98-D: MAb peptide reactivity pattern clustered with immunological related MAbs: 1108, 386, 268, 311, 257, 694.8
   the amino acids HI tended to be critical for reactivity in this group. Zolla-Pazner *et al.* [1999b]
- 694/98-D: Used to study pre- and post-exposure prophylaxis Hu-PBL-SCID mice infected by an intraperitoneal injection of HIV-1 LAI MAb half-life in plasma in mice is 9 days 2 hours post-694/98-D mice were challenged with LAI, and at an Ab concentration of 1.32 mg/Kg, 50% of the mice were infected one of the infected mice carried the resistant form GRTF rather than GRAF (critical amino acids for binding are GRA) post-exposure prophylaxis was effective if delivered 15 min post-exposure, but declined to 50% if delivered 60 min post-exposure, and similar time constraints have been observed for HIVIG, 2F5 and 2G12, in contrast to MAb BAT123 that could protect delivered 4 hours post infection. Andrus *et al.* [1998]
- 694/98-D: Neutralization synergy was observed when the MAbs 694/98-D (V3), 2F5 (gp41), and 2G12 (gp120 discontinuous) were used in combination, and even greater neutralizing potential was seen with the addition of a fourth MAb, F105 (CD4 BS) Li *et al.* [1998]
- 694/98-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H – 694/98-D bound only to B and D clade virions and had limited cross reactivity. Nyambi *et al.* [1998]
- 694/98-D: In a study of the influence of the glycan at position 306 of the V3 loop on MAb recognition, anti-V3 MAbs were found to neutralize an HIV-BRU mutant virus that lacks the V3 loop glycan more efficiently than HIV-BRU. Schonning *et al.* [1998]
- 694/98-D: The tip of the MN V3 loop was inserted into cold causing human rhinovirus 14 (HRV14) chimeras were immunoselected, and chimeric viruses were neutralized by anti-V3 loop antibodies, and 694/98-D was among the Abs used chimeric viruses elicited potent NAbs in guinea pigs against ALA-1 and MN. Smith et al. [1998]

- 694/98-D: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env could only achieve 50% neutralization alone all Ab combinations tested showed synergistic neutralization 694/98-D has synergistic response with MAbs F105, 15e, b12, 2F5, 17b, 2G12, and 48d, and with HIVIG. Li *et al.* [1997]
- 694/98-D: ADCC activity, and no viral enhancing activity.
   Forthal et al. [1995]
- 694/98-D: Human HIV-1 infected sera and MAb 694/98 have high reactivity to MN and RF infected H9 cells, but Genentech rec gp120 IIIB vaccine recipients do not. VanCott et al. [1995]
- 694/98-D: Serotyping study using flow-cytometry bound GRAX bearing virus in 10/11 cases – somewhat conformation dependent. Zolla-Pazner et al. [1995]
- 694/98-D: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon V3 MAbs can inhibit gp120 binding to GalCer *in vitro* binding of GalCer to gp120 inhibited but did not completely block MAb binding. Cook *et al.* [1994]
- 694/98-D: 50% neutralization of HIV-IIIB at a concentration of 0.15mug/ml. Gorny et al. [1994]
- 694/98-D: Potent neutralization of IIIB no neutralization synergy in combination with CD4 binding domain MAbs. Laal *et al.* [1994]
- 694/98-D: GRVY did not alter peptide binding GRVI and GQAW enhanced dissociation – GQVF and GQAL did not bind. VanCott et al. [1994]
- 694/98-D: Neutralizes MN and IIIB (GRAF) binds SF2 (GRAF) binding reactivity: MN, IIIB, SF2, NY5, RF, CDC4, WM52. Gorny *et al.* [1993]
- 694/98-D: Called 694-D complement mediated virolysis of IIIB, but not in the presence of sCD4. Spear *et al.* [1993]
- 694/98-D: Type-specific lab isolate neutralization was observed

   binds with 1-3 fold greater affinity to gp120 than to peptides.
   Gorny et al. [1992]
- 694/98-D: This MAb was first described here. Skinner *et al.* [1988b]

**No.** 563

**MAb ID** MO101/V3,C4

**HXB2 Location** gp160 (314–323)

**Author Location** gp120 (314–323)

Epitope GRAFVTIGKI+LGVAPTKAKR

**Neutralizing** 

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgM)

Ab Type gp120 V3-C4

References Ohlin et al. 1992

MO101: Generated in response to IIIB Env 286-467 upon in vitro stimulation of uninfected-donor lymphocytes – reacts with peptides 314-323 + 494-503 from the V3 and C4 regions. Ohlin et al. [1992]

No. 564

**MAb ID** MO101/V3,C4

HXB2 Location gp160 (314-323)

Author Location gp120 (314–323)

Epitope GRAFVTIGKI+LGVAPTKAKR

Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgM) **Ab Type** gp120 V3-C5 References Ohlin et al. 1992

• MO101: Generated through in vitro stimulation of uninfecteddonor lymphocytes with pB1 containing IIIB Env 286-467 reacts with peptides from the V3 and C4 regions, positions 314-323 + 494-503, peptides GRAFVTIGKI + LGVAPTKAKR. Ohlin et al. [1992]

No. 565

MAb ID MO101/V3,C4

HXB2 Location gp160 (314-323)

Author Location gp120 (494-503)

Epitope GRAFVTIGKI+LGVAPTKAKR

**Neutralizing** 

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgM)

**Ab Type** gp120 V3-C5

References Ohlin et al. 1992

• MO101: Generated through in vitro stimulation of uninfected- Species (Isotype) mouse donor lymphocytes with pB1 containing IIIB Env 286-467 reacts with peptides from the V3 and C4 regions, positions 314- Research Contact F. Traincard, Pasteur Institute, France 323 + 494-503, peptides GRAFVTIGKI + LGVAPTKAKR. Ohlin et al. [1992]

**No.** 566

**MAb ID** 9205 (NEA-9205 NEA9205)

**HXB2 Location** gp160 (315–317)

Author Location gp120 (IIIB)

Epitope RAF(corereactivity)

Neutralizing L

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

HIV component: V3

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

Research Contact NEN, Boston MA, commercial

References Gram et al. 2002; Schonning et al. 1999; Schonning et al. 1998; Fontenot et al. 1995; VanCott et al. 1994; Allaway et al. 1993; Tru-

jillo et al. 1993; Durda et al. 1990

• 9205: Also see MAb called 5023A.

- 9205: Called NEA9205 gp120 capture ELISAs with MAbs D7324 (anti-C-term) or 9205 (anti-V3) were compared in a study of orientation of glycosylation sites - CD4 binding could only inhibit deglycosylation when gp120 was bound to the plate by D7324, not by 9205, while Abs from HIV-1 infected people inhibited deglycosylation most effectively when gp120 was caught by 9205. Gram et al. [2002]
- 9205: Called NEA-9205 the stoichiometry of MAb neutralization was tested and the data indicated that binding for neutralization was was incremental not all or none, i.e., each envelope oligomer binds a single MAb and each Env oligomer bound reduces the chances of infection - 9205 binds only to Env with a glycosylation site mutation in the V3 loop, A308T. Schonning et al. [1999]
- 9205: Called NEA-9205 The N306 glycan of the V3 loop makes the tip of the V3 loop inaccessible to this MAb in oligomeric Env, loss of this glycan enhances neutralization sensitivity. Schonning et al. [1998]

• 9205: Neutralizes IIIB but not MN – significantly slower dissociation constant for IIIB than MN. VanCott et al. [1994]

- 9205: Synergy with combinations of CD4-based molecules in inhibition of HIV-1 Env mediated cell fusion. Allaway et al. [1993]
- 9205: Called NEA-9205, epitope RIQRGPGRAFVTIGK reacts with three human brain proteins of 35, 55, 110 kd molecular weight – similar to 9284 – RAF is the core reactivity. Trujillo et al. [1993]

No. 567

**MAb ID** 110.I

HXB2 Location gp160 (316-322)

**Author Location** gp120 (316–322)

Epitope AFVTIGK

Neutralizing L

Immunogen vaccine

Vector/Type: protein HIV component:

gp120

Ab Type gp120 V3

References Parren et al. 1998a; Wyatt et al. 1997; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; Moore et al. 1994c; Moore et al. 1993b

- 110.I: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated - authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- 110.I: Binds both gp120 and soluble gp120+gp41 complex efficiently, suggesting its gp120 epitope is not blocked by gp41 binding. Wyatt et al. [1997]
- 110.I: Reciprocal binding inhibition with other anti-V3 and anti-C4 MAbs - and enhances binding of some anti-V2 MAbs binding enhanced by some anti-CD4 binding site MAbs. Moore & Sodroski [1996]
- 110.I: Epitope suggested to be RAFVTIGK V3 MAbs 9284, BAT123, 110.5, and 110.I could each significantly increase gp120 dissociation from virus, mimicking sCD4, and expose the gp41 epitope for MAb 50-69, in contrast to anti-V2 MAbs. Poignard et al. [1996a]
- 110.I: Binds equally well to monomer and oligomer, rapid association and potent neutralization of lab strains. Sattentau & Moore [1995]
- 110.I: Binds to carboxy-terminal side of the V3 loop inhibits binding of C4 region MAb G3-299. Moore et al. [1993b]

No. 568

MAb ID anti-HIV-2 polyclonal

HXB2 Location gp160 (317–320)

Author Location gp120 (315-318 SBL6669 HIV-2)

Epitope FHSQ+WCR

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide Strain: HIV-2

SBL6669-ISY HIV component: V3

Species (Isotype) guinea pig (IgG)

Ab Type gp120 V3

References Morner et al. 1999

• Neutralizing Abs against HIV-2 V3 are produced when peptides spanning two non-contiguous parts of the V3 loop are used for vaccination including amino acids 315-318 near the tip (FHSQ) and 329-331 (WCR) at the C-term Cys. Morner *et al.* [1999]

No. 569

MAb ID IIIB-V3-01

HXB2 Location gp160 (320-328)

Author Location gp120 (IIIB)

Epitope IGKIGNMRQ

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

HIV component: V3

Species (Isotype) mouse (IgG1)

Ab Type gp120 V3

Research Contact Jon Laman

References Laman et al. 1993

- IIIB-V3-01: UK Medical Research Council AIDS reagent: ARP3046
- IIIB-V3-01: NIH AIDS Research and Reference Reagent Program: 1726.
- IIIB-V3-01: Specific for carboxy-terminal flank of the IIIB V3 loop – epitope is hidden native gp120, exposed on denaturation. Laman et al. [1993]

**No.** 570

MAb ID D/6D1

**HXB2 Location** gp160 (346–377)

Author Location gp120 (351–382 LAI)

Epitope ASKLREQFGNNKTIIFKQSSGGDPEIVTHSFN

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 V4

References Bristow et al. 1994

D/6D1: V4 MAb generated in a study of the humoral immune response to Baculovirus-expressed mis-folded rgp120 and rgp160. Bristow et al. [1994]

**No.** 571

**MAb ID** 4D7/4

**HXB2 Location** gp160 (360–380)

Author Location gp120 (361–380 LAI)

 ${\bf Epitope} \ \, {\tt IFKQSSGGDPEIVTHSFNCGG}$ 

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 V4

Research Contact S. Ranjbar, NIBSC, UK

References Moore et al. 1994c

 4D7/4: UK Medical Research Council AIDS reagent: ARP3051.

• 4D7/4: C3 region – the relative affinity for denatured/native gp120 is >10. Moore *et al.* [1994c]

**No.** 572

**MAb ID** 36.1(ARP 329)

**HXB2 Location** gp160 (361–381)

Author Location gp120 (362-381 LAI)

Epitope FKQSSGGDPEIVTHSFNCGGE

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 V4

References Moore et al. 1994c; Thiriart et al. 1989

- 36.1: UK Medical Research Council AIDS reagent: ARP329.
- 36.1: The relative affinity for denatured/native gp120 is >30 mutations 380 G/F, 381 E/P impair binding. Moore *et al.* [1994c]

**No.** 573

MAb ID C12

HXB2 Location gp160 (361–381)

Author Location gp120 (362-381 LAI)

Epitope FKQSSGGDPEIVTHSFNCGGE

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG1)

Ab Type gp120 V4

Research Contact George Lewis

**References** Moore *et al.* 1994d; Abacioglu *et al.* 1994; Moore *et al.* 1994c; Moore & Ho 1993

- C12: C3 region epitope boundaries mapped by peptide scanning, core FNCGG. Abacioglu *et al.* [1994]
- C12: The relative affinity for denatured/native gp120 is >30 mutations 380 G/F, 381 E/P, and 384 Y/E impair binding also binds GEFFYCNSTQLFNS, gp120(380-393 LAI) Moore *et al.*
- C12: Bound preferentially to denatured IIIB gp120. Moore & Ho [1993]

**No.** 574

**MAb ID** 110.D

HXB2 Location gp160 (380-393)

Author Location gp120 (380–393 LAI)

**Epitope** GEFFYCNSTQLFNS

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: Env

Species (Isotype) mouse (IgG)

**Ab Type** gp120 C3

Research Contact F. Traincard, Pasteur Institute, France

References Valenzuela et al. 1998; Moore et al. 1994c

• 110.D: The relative affinity for denatured/native gp120 is >50. Moore *et al.* [1994c]

No. 575

MAb ID B32

**HXB2 Location** gp160 (380–393)

**Author Location** gp120 (380–393 LAI) **Epitope** GEFFYCNSTQLFNS

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG1)

Ab Type gp120 C3

References Abacioglu et al. 1994; Moore et al. 1994c

- B32: C3 region epitope boundaries mapped by peptide scanning FFY(core) Abacioglu et al. [1994]
- B32: The relative affinity for denatured/native gp120 is >100 mutations 380 G/F, 381 G/P, 382 F/L, 384 Y/E, and 386 N/R impair binding. Moore *et al.* [1994c]

**No.** 576

MAb ID polyclonal (VEI4)

HXB2 Location gp160 (391–413)

Author Location Env

**Epitope** FNSTWFNSTWSTEGSNNTEGSDT

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V4

References Carlos et al. 1999

Antibody response to the epitopes in a vaccine construct (VEI) containing peptides from 5 hypervariable regions of gp120 was detected in the sera of HIV-1 positive subjects, including sera from 6 non-subtype B infections – serum samples from San Francisco, Canada and Puerto Rico cohort showed presence of antibodies against all five VEI hypervariable regions, but most consistently against the V3 region peptide NNNTRKSIRIGP-GRAFYTTGDIGNIRQ. Carlos et al. [1999]

No. 577

MAb ID B15

**HXB2 Location** gp160 (395–400)

**Author Location** gp120 (395–400 BH10)

**Epitope** WFNSTW

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG2b)

Ab Type gp120 V4

Research Contact George Lewis

**References** Abacioglu *et al.* 1994; Moore *et al.* 1993b; Moore & Ho 1993

- B15: V4 region epitope boundaries mapped by peptide scanning. Abacioglu et al. [1994]
- B15: Bound preferentially to denatured IIIB gp120. Moore & Ho [1993]

 B15: Binds native BH10 gp120 with 5 fold less affinity than denatured – does not bind native or denatured MN gp120. Moore et al. [1993b]

**No.** 578

MAb ID B34

HXB2 Location gp160 (395–400)

**Author Location** gp120 (395–400 BH10)

**Epitope** WFNSTW

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG2b)

Ab Type gp120 V4

References Abacioglu et al. 1994

B34: V4 region – epitope boundaries mapped by peptide scanning. Abacioglu et al. [1994]

**No.** 579

MAb ID 7F11

**HXB2 Location** gp160 (397–439)

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component:

gp120

Species (Isotype) mouse

References Nilsen et al. 1996; Lasky et al. 1987

• 7F11: There is another MAb with this name that binds to integrase. Nilsen *et al.* [1996]

**No.** 580

MAb ID E51

**HXB2 Location** gp160 (420–423)

Author Location gp120 (420–423 HXB2)

Epitope IKQI

Subtype B

Neutralizing P

Immunogen

Species (Isotype) human

Ab Type gp120 CD4i, gp120 CCR5BS

Research Contact Joseph

joseph\_sodroski@dfci.harvard.edu

References Haynes et al. 2005; Xiang et al. 2003

Keywords antibody binding site definition and expo-

sure, antibody generation, co-receptor, subtype comparisons, variant cross-recognition

Sodroski,

or cross-neutralization

• E51: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. E51 has no indication of polyspecific autoreactivity. Haynes *et al.* [2005] (antibody binding site definition and exposure)

• E51: E51 recognizes a highly conserved epitope localized in the basic  $\beta$ 19-strand (gp120 aa420-423), a region involved in B-cell line established from an HIV+ individual undergoing early STI. Fab fragments were also produced. E51, like CD4i MAb 17b, blocks CCR5 binding to sCD4-bound gp120. The presence of sCD4 induces a conformational change in gp120, which enhances ligand recognition. The substitutions E381R, F383S, R419D I420R, K421D, Q422L, I423S, and Y435S (HXB2 numbering) all severely reduce 17b and E51 binding. All but I423S also diminish CCR5 binding by more than 50%. The mutation F383S also inhibits sCD4 binding and CD4BS MAb F105 binding, and K421D inhibits F105 binding, but not sCD4. E51 has more cross-neutralizing potency than other prototype CD4i MAbs (17b) for B and C clade isolates. E51 and 17b both neutralized HIV-1 clade B strains HXBc2 and ADA, while JR-FL and 89.6 were only neutralized by E51, not 17b. Clade C strains MCGP1.3 and SA32 were both inhibited by 17b and E51, but E51 was more potent against SA32. Xiang et al. [2003] (antibody binding site definition and exposure, antibody generation, co-receptor, variant cross-recognition or cross-neutralization, subtype comparisons)

No. 581

MAb ID JL413

HXB2 Location gp160 (421-436)

Author Location gp160 (421-436)

Epitope KQIINMWQEVGKAMYA

Subtype B Neutralizing P

Immunogen autoimmune disease

Species (Isotype) human

**Ab Type** gp120 CD4BS **References** Karle *et al.* 2004

**Keywords** antibody generation, antibody sequence, variable domain, co-receptor, subtype compariable

sons

• JL413: Phage display was used to create a library of gp120-binding single-chain fragments containing V domain (scFv) constructs derived from PBMC of lupus patients. Lupus patients rarely get HIV/AIDS and can make antibodies that bind to a conserved gp120 determinant. The scFV clone JL413 was able to induce dose-dependent, cross-clade neutralization of primary HIV-1 isolates ZA009 (R5, clade C); BR004 (R5, clade C); Ug046 (X4, clade D); SF162 (R5, clade B), and 231135 (clade B). The scFV clone JL413 recognizes a linear region that overlaps the CD4 T-cell binding site, in contrast to HIV-induced MAbs that bind to this region and are conformation dependent. Karle et al. [2004] (antibody generation, co-receptor, subtype comparisons, antibody sequence, variable domain)

**No.** 582

MAb ID 5C2E5

HXB2 Location gp160 (422–431)

Author Location gp120 (406–415 IIIB)

Epitope QFINMWQEVK

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component:

gp120

Species (Isotype) mouse

Ab Type gp120 C4

CCR5 binding. The MAb was isolated from a EBV transformed Research Contact T. Gregory and R. Ward, Genentech, San Fran-

cisco

References Cordell et al. 1991; Lasky et al. 1987

• 5C2E5: Cross-competition with MAbs 5C2E5, ICR38.8f and ICR38.1a. Cordell *et al.* [1991]

• 5C2E5: Blocks the gp120-CD4 interaction. Lasky et al. [1987]

**No.** 583

**MAb ID** G3-211 (211/C)

**HXB2 Location** gp160 (423–437)

Author Location gp120 (423-437 IIIB)

Epitope IINMWQKVGKAMYAP

**Neutralizing** L

Immunogen vaccine

Vector/Type: virus derived protein Strain: B clade IIIB HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 C4

References Pantophlet et al. 2004; Sun et al. 1989

Keywords vaccine antigen design

- G3-211: Note it is not clear if 211/C and G3-211 are the same antibodies. By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including 211/C Pantophlet *et al.* [2004] (vaccine antigen design)
- G3-211: G3-211, 42, 299, 508, 519, 536, 537: Cross-react with diverse strains by immunofluorescence blocks HIV binding to CD4+ cells different neutralization efficiencies. Sun *et al.* [1989]

No. 584

**MAb ID** G3-537

HXB2 Location gp160 (423-437)

**Author Location** gp120 (423–437 IIIB)

Epitope IINMWQKVGKAMYAP

**Neutralizing** L

Immunogen vaccine

Vector/Type: virus derived protein Strain: B

clade IIIB HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 C4

**References** Zwick *et al.* 2003; McKeating *et al.* 1992b; Ho *et al.* 1991b; Sun *et al.* 1989

**Keywords** antibody interactions

G3-537: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact

4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the C4 MAbs used. Zwick *et al.* [2003] (antibody interactions)

- G3-537: Weakly neutralizing binds to a linear binding domain of gp120, NMWQEVGKAMYAPPISG. McKeating *et al.* [1992b]
- G3-537, 211, 299, 508, 519, 536, 42: Cross-react with diverse strains by immunofluorescence – blocks HIV binding to CD4+ cells – different neutralization efficiencies. Sun *et al.* [1989]

No. 585

MAb ID polyclonal

HXB2 Location gp160 (425-436)

Author Location gp120

Epitope NMWQEVGKAMYA

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: B clade IIIB

Adjuvant: Cholera toxin (CT)

Species (Isotype) mouse (IgA)

Ab Type gp120 CD4BS

References Bukawa et al. 1995

Polyclonal secretory IgA antibody raised by mucosal immunization is able to neutralize IIIB, SF2, and MN – HIV-1 neutralization may be due to the V3, CD4 or HPG30 component of the multicomponent peptide immunogen. Bukawa et al. [1995]

**No.** 586

**MAb ID** 1795

HXB2 Location gp160 (425-441)

**Author Location** gp120 (425–441 IIIB)

Epitope NMWQEVGKAMYAPPISG

Neutralizing L

Immunogen vaccine

Vector/Type: poliovirus HIV component:

Env

Species (Isotype)

Ab Type gp120 CD4BS

References McKeating et al. 1992b

1795: CD4 binding site – weakly neutralizing – binding inhibited by WQEVGKAMYA, GKAM may be involved. McKeating et al. [1992b]

**No.** 587

**MAb ID** ICR38.1a (38.1a, 388/389, ARP388/389)

**HXB2 Location** gp160 (429–438)

**Author Location** gp120 (427–436 BRU)

Epitope EVGKAMYAPP

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2b)

**Ab Type** gp120 C3, gp120 C4

References Vella et al. 2002; Kropelin et al. 1998; Peet et al. 1998; Jeffs et al. 1996; Moore et al. 1993b; McKeating et al. 1993a; McKeating et al. 1992c; McKeating et al. 1992a; McKeating et al. 1992b; Cordell et al. 1991

- ICR38.1a: UK Medical Research Council AIDS reagent: ARP388/ARP389.
- ICR38.1a: Called ARP388/ARP389: Herpesvirus saimiriimmortalized CD4+ T lymphocytes (HVS T cells) were used to isolate virus and perform HIV-1 neutralization assays, and compared with a standard PBMC protocol – neutralization sensitivities to a panel of MAbs and to homologous or heterologous plasma/sera were similar for HVS T cells (CN-2 cells) and PBMCs – lists epitope as WQEVGKAMYA. Vella *et al.* [2002]
- ICR38.1a: Called 388/389 anti-C1 region MAb 87-135/9 blocks gp120 interaction with CD4+ cells blocking activity is additive when combined with antibodies which bind in the C4 region of gp120 (F105, 388/389, and b12) Kropelin *et al.* [1998]
- ICR38.1a: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind ICR38.1a was not affected by V3 serine substitutions mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet *et al.* [1998]
- ICR38.1a: Called 38.1a 10 to 20 fold increased binding when V1/V2 or V1/V2 and V3 were deleted from gp120. Jeffs *et al.* [1996]
- ICR38.1a: Studied in the context of a neutralization escape mutant. McKeating *et al.* [1993a]
- ICR38.1a: Unreactive with solid-phase decapeptide, competed in solution phase assay ICR 38.1a and ICR 38.8f were initially reported to be independent MAbs, but are actually subclones of the same MAb. Moore *et al.* [1993b]
- ICR38.1a: Weakly neutralizing binds linear determinant in the CD4 binding domain cross-competition with MAbs G3-536, 5C2E5, and ICR38.8f. Cordell *et al.* [1991]; McKeating *et al.* [1992b]
- ICR38.1a: Unable to exert a synergistic effect in combination with V3 directed MAbs, in contrast to MAb 39.13g, that binds to a conformational epitope involved in CD4 binding. McKeating et al. [1992a]

No. 588

MAb ID G3-299

HXB2 Location gp160 (429–438)

**Author Location** gp120 (429–438 BRU)

Epitope EVGKAMYAPP

**Neutralizing** L

Immunogen vaccine

Vector/Type: virus derived protein HIV com-

ponent: gp120

Species (Isotype) mouse (IgG1)

**Ab Type** gp120 C4

**Research Contact** M. Fung and Tanox Biosystems Inc and David

Ho, ADARC, NY

ren et al. 1998a; Wyatt et al. 1997; Ditzel et al. 1997; Binley et al. 1997a; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; Moore et al. 1993b; Sun et al. 1989

**Keywords** antibody binding site definition and exposure, antibody interactions

- G3-299: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the C4-V3 MAbs used. Zwick et al. [2003] (antibody interactions)
- G3-299: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-299) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Authors describe the G3-299 epitope as V3 loop/outer domain. Kwong et al. [2002] (antibody binding site definition and exposure)
- G3-299: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated – authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- G3-299: Binds both gp120 and soluble gp120+gp41 complex efficiently, suggesting its gp120 epitope is not blocked by gp41 binding. Wyatt et al. [1997]
- G3-299: Discontinuous V3-C4 epitope, binding enhanced by a few anti-C1, anti-CD4 binding site, and V2 MAbs - binding reciprocally inhibited by anti-V3 MAbs - G3-229 enhances the binding of some anti-V2 MAbs. Moore & Sodroski [1996]
- G3-299: Epitope described as KQIINMWQKVGKAMYAPPIS - binding resulted in slight gp120 dissociation from virus and exposure of the gp41 epitope for MAb 50-69. Poignard et al. [1996a]

- References Zwick et al. 2003; Kwong et al. 2002; Par- G3-299: Binds with higher affinity to monomer than to oligomer, slow association rate, although faster than other C4 MAbs tested, with more potent neutralization of lab strain. Sattentau & Moore [1995]
  - G3-299: C4 region - binds HXB2 20mer KQIIN-MWQKVGKAMYAPPIS, and SF-2 and MN gp120s - G3-42, G3-299 lower affinity than G3-508, G3-519, and G3-536 - bound native gp120, not denatured - poor peptide binding, epitope spans V3-C4 regions – 433A/L, 435Y/H and 430V/S substitutions impaired binding, V3 loop cleavage or insertion abolished binding. Moore et al. [1993b]
  - G3-299: Best neutralization of IIIB in panel of 7 MAbs that bind overlapping epitope. Sun et al. [1989]

No. 589

**MAb ID** G3-42 (G3 42)

HXB2 Location gp160 (429-438)

Author Location gp120 (429–438 BRU)

Epitope EVGKAMYAPP

**Neutralizing** L

Immunogen vaccine

Vector/Type: virus derived protein Strain: B

clade IIIB HIV component: gp120

**Species (Isotype)** mouse (IgG1)

Ab Type gp120 C4

Research Contact Tanox Biosystems Inc and David Ho, ADARC, NY, NY

References Koefoed et al. 2005; Zwick et al. 2003; Jagodzinski & Trzeciak 2000; Binley et al. 1999; Binley et al. 1997a; Trkola et al. 1996a; Poignard et al. 1996a; Moore & Sodroski 1996; Jagodzinski et al. 1996; Sattentau & Moore 1995; Thali et al. 1993; Moore et al. 1993b; Sun et al. 1989

Keywords antibody binding site definition and exposure, antibody interactions

- G3-42: IgG antibody phage display libraries were created from HIV-1 + individuals after pre-selection of PBMC with gp120, as an alternative to using bone marrow for generating libraries. G3-42 was among a set of Abs used for competition studies to define the binding sites of the newly isolated MAbs, representing a MAb with a linear C4/V3 epitope. Koefoed et al. [2005] (antibody binding site definition and exposure)
- G3-42: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the C4-V3 MAbs used. Zwick et al. [2003] (antibody interactions)
- 0.5beta: MAbs 0.5beta and G3-42 were used to study synthesis of oligomeric and monomeric forms of Env - inhibition of glycosylation by tunicamycin results in reduction of oligomeric gp120 at the cell surface and of monomer in the

cytoplasm - neither MAb recognized non-glycosylated Env precursor. Jagodzinski & Trzeciak [2000]

- G3-42: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits – a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen – SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 - nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999]
- G3-42: The sulfated polysaccharide curdlan sulfate (CRDS) binds to the Envelope of T-tropic viruses and neutralizes virus - CRDS potently inhibits G3-42 binding - G3-42 epitope described as KVGKAMYAPP. Jagodzinski et al. [1996]
- G3-42: Inhibits binding of many anti-V3, -CD4 binding site, and -C4 region MAbs - enhances binding of some anti-V2 region MAbs. Moore & Sodroski [1996]
- G3-42: Epitope described as KQIINMWQKVGKAMYAPPIS - binding resulted in slight gp120 dissociation from virus and exposure of the gp41 epitope for MAb 50-69. Poignard et al. [1996a]
- G3-42: Called G3 42 Does not inhibit gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study – described as V3-C4 discontinuous epitope. Trkola et al. [1996a]
- G3-42: Binds with higher affinity to monomer than to oligomer, slow association rate. Sattentau & Moore [1995]
- G3-42: C4 region - binds HXB2 20mer KQIIN-MWQKVGKAMYAPPIS, and SF-2 and MN gp120s - G3-42, G3-299 have lower affinity than G3-508, G3-519, and G3-536 - bound native gp120, not denatured - poor peptide binding, epitope spans V3-C4 regions - 433A/L, 435Y/H and 430V/S substitutions impaired binding, V3 loop insertion abolished binding. Moore et al. [1993b]
- G3-42: Inhibits binding of CD4 inducible MAb 48d. Thali et al. [1993]
- G3-42: Neutralization of IIIB but not RF. Sun et al. [1989]

**No.** 590

MAb ID G3-508 (G3 508)

HXB2 Location gp160 (429-438)

Author Location gp120 (429–438 BRU)

Epitope EVGKAMYAPP

Neutralizing L

Immunogen vaccine

Vector/Type: virus derived protein Strain: B clade IIIB HIV component: gp120

**Species (Isotype)** mouse (IgG1)

Ab Type gp120 C4

**Research Contact** M. Fung and Tanox Biosystems Inc and David Ho, ADARC, NY

References Binley et al. 1998; Parren et al. 1998a; Binley et al. 1997a; Trkola et al. 1996a; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; Moore et al. 1993b; Thali et al. 1993; Sun et al. 1989

- G3-508: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein (Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer. Binley et al. [1998]
- G3-508: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated – authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- G3-508: Inhibits binding of some V3, C4 and CD4 binding site MAbs, enhances binding of V2 region MAbs. Moore & Sodroski [1996]
- G3-508: Binding resulted in slight gp120 dissociation from virus and exposure of the gp41 epitope for MAb 50-69. Poignard et al. [1996a]
- G3-508: Called G3 508 inhibits gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]
- G3-508: Binds with higher affinity to monomer than to oligomer, slow association rate. Sattentau & Moore [1995]
- C4 region binds HXB2 20mer KQIIN-MWQKVGKAMYAPPIS, and SF-2 and MN gp120s - bound denatured with 10 fold greater affinity than native - 433A/L, 435Y/H and 430V/S substitutions impaired binding. Moore et al. [1993b]
- G3-508: Inhibits binding of CD4 inducible MAb 48d. Thali et al. [1993]
- G3-508: Neutralization of IIIB and RF. Sun et al. [1989]

No. 591

MAb ID G3-519

HXB2 Location gp160 (429–438)

Author Location gp120 (429-438 BRU)

**Epitope** EVGKAMYAPP

**Neutralizing** L

Immunogen vaccine

Vector/Type: virus derived protein Strain: B clade IIIB HIV component: gp120

**Species (Isotype)** mouse (IgG1)

Ab Type gp120 C4

Research Contact Tanox Biosystems Inc and David Ho,

ADARC, NY

References Zwick et al. 2003; Binley et al. 1999; Parren et al. 1998a; Wyatt et al. 1997; Binley et al. 1997a; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; D'Souza et al. 1994: Moore et al. 1993b: Moore & Ho 1993; Sun et al. 1989

Keywords antibody interactions

• G3-519: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding

to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the C4 MAbs used. Zwick et al. [2003] (antibody interactions)

- G3-519: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits – a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen – SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 – nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999]
- G3-519: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated – authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- G3-519: Binds both gp120 and soluble gp120+gp41 complex efficiently, suggesting its gp120 epitope is not blocked by gp41 binding. Wyatt et al. [1997]
- G3-519: Non-reciprocal enhanced binding in the presence of the C5 MAb 1C1 and the C1 MAb 135/9 - reciprocal enhanced binding with some V2 MAbs. Inhibited binding the presence of some C4, V3 and CD4 binding site MAbs. Moore & Sodroski [1996]
- G3-519: Epitope described as KVGKAMYAPP binding resulted in slight gp120 dissociation from virus but no significant exposure of the gp41 epitope for MAb 50-69. Poignard et al. [1996a]
- G3-519: Binds with higher affinity to monomer than to oligomer, slow association rate. Sattentau & Moore [1995]
- G3-519: Included in a multi-lab study for antibody characterization, and binding and neutralization assay comparison, also binds IIIB: IINMWQKVGKAMYAPP. D'Souza et al. [1994]
- G3-519: Neutralizes IIIB, is reactive with SF-2 gp120, mild inhibition of HIV-1 + sera binding to IIIB gp120. Moore & Ho [1993]
- C4 region binds HXB2 20mer KQIIN-• G3-519: MWQKVGKAMYAPPIS, and SF-2 and MN gp120s - bound denatured with 5 fold greater affinity than native - 433A/L, 435Y/H, 438P/R and 430V/S substitutions impaired binding. Moore et al. [1993b]
- G3-519: Best neutralization of RF in panel of 7 MAbs that bind overlapping epitope. Sun et al. [1989]

No. 592

**MAb ID** G3-536

**HXB2 Location** gp160 (429–438)

Author Location gp120 (429–438 BRU)

Epitope EVGKAMYAPP

**Neutralizing** L

Immunogen vaccine

Vector/Type: virus derived protein Strain: B

clade IIIB HIV component: gp120

**Species (Isotype)** mouse (IgG1)

Ab Type gp120 C4

Research Contact Tanox Biosystems Inc and David Ho,

ADARC, NY

References Parren et al. 1998a; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; Gorny et al. 1994; Moore et al. 1993b; Moore & Ho 1993; McKeating et al. 1992b; Cordell et al. 1991; Ho et al. 1991b; Sun et al.

- G3-536: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated – authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]
- G3-536: Inhibits binding of some V3, C4 and CD4 binding site MAbs, enhances binding of V2 region MAbs. Moore & Sodroski [1996]
- G3-536: Epitope described as KVGKAMYAPP. Poignard et al. [1996a]
- G3-536: Binds with higher affinity to monomer than to oligomer, slow association rate. Sattentau & Moore [1995]
- G3-536: Enhances binding of anti-V2 MAb 697-D. Gorny et al. [1994]
- G3-536: Neutralizes IIIB, is reactive with SF-2 gp120, mild inhibition of HIV-1 + sera binding to IIIB gp120. Moore & Ho [1993]
- G3-536: C4 region - binds HXB2 20mer KQIIN-MWQKVGKAMYAPPIS, and SF-2 and MN gp120s - bound denatured with 15 fold greater affinity than native – 433A/L, 435Y/H, 438P/R, and 430V/S substitutions impaired binding. Moore et al. [1993b]
- G3-536: Weakly neutralizing binds to a linear determinant in the CD4 binding domain of gp120. McKeating et al. [1992b]
- G3-536: Cross-competition with MAbs 5C2E5, ICR38.8f and ICR38.1a. Cordell et al. [1991]
- G3-536: Weak neutralization of IIIB and RF cross-react with diverse strains by immunofluorescence - blocks HIV binding to CD4+ cells - epitope: IINMWQKVGKAMYAP. Sun et al. [1989]

No. 593

MAb ID ICR38.8f

**HXB2 Location** gp160 (429–438)

Author Location gp120 (429–438 BRU)

Epitope EVGKAMYAPP

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2b)

Ab Type gp120 C4

References Moore et al. 1993b; Cordell et al. 1991

• ICR38.8f:ICR 38.1a and ICR 38.8f were initially reported to be independent MAbs, but are actually subclones of the same MAb. Moore *et al.* [1993b]

• ICR38.8f: Weakly neutralizing – binds linear determinant in the CD4 binding domain – cross-competition with ICR38.1a, 5C2E5, and G3-536. Cordell *et al.* [1991]

**No.** 594

MAb ID MO86/C3

HXB2 Location gp160 (429–443)

**Author Location** gp120 (429-443)

Epitope EVGKAMYAPPISGQI

Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgM)

Ab Type gp120 C4

References Ohlin et al. 1992

 MO86: Generated in response to IIIB Env 286-467 upon in vitro stimulation of uninfected-donor lymphocytes. Ohlin et al. [1992]

No. 595

**MAb ID** 13H8

**HXB2 Location** gp160 (431–440)

**Author Location** gp120 (412–453)

**Epitope** GKAMYAPPIS

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade MN

 $\boldsymbol{Species}\;(\boldsymbol{Isotype})\;\;\text{mouse}\;(\boldsymbol{IgG})$ 

Ab Type gp120 C4

**References** Jeffs *et al.* 1996; Nakamura *et al.* 1993; Nakamura *et al.* 1992

- 13H8: Binds V3 and C4 peptides (J. P. Moore, per. comm.)
- 13H8: 3 and 4.5 fold increased binding when V1/V2 or V1/V2 and V3 were deleted from gp120, respectively. Jeffs *et al.* [1996]
- 13H8: Bound diverse strains, neutralizing activity against MN. Nakamura *et al.* [1993]
- 13H8: Cross blocks 5C2 in IIIB-rsgp160 ELISA reactive with diverse strains in rgp120 ELISA. Nakamura *et al.* [1992]

No. 596

**MAb ID** G45-60

HXB2 Location gp160 (431-440)

Author Location gp120 (429–438 BRU)

Epitope GKAMYAPPIS

Neutralizing L

Immunogen vaccine

Vector/Type: virus derived protein Strain: B clade IIIB HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 C4

**References** Jagodzinski *et al.* 1996; Moore & Sodroski 1996; Gorny *et al.* 1994; Moore *et al.* 1993b;

Sun et al. 1989

• G45-60: The sulfated polysaccharide curdlan sulfate (CRDS) binds to the Envelope of T-tropic viruses and neutralizes virus CRDS inhibits G45-60 binding. Jagodzinski *et al.* [1996]

 G45-60: Non-reciprocal enhancement of G45-60 binding by some C1 and C5 antibodies – reciprocal enhancement of some V2 region MAbs – reciprocal inhibition with many MAbs that bind to the V3, C4 and CD4 binding site regions. Moore & Sodroski [1996]

- G45-60: Enhances binding of anti-V2 MAb 697-D. Gorny et al. [1994]
- G45-60: C4 region binds HXB2 20mer KQIIN-MWQKVGKAMYAPPI, decapeptide flanking peptides also bound bound equivalently to native and denatured gp120 433A/L and 435Y/H (not 430V/S) substitutions impaired binding. Moore *et al.* [1993b]

**No.** 597

MAb ID polyclonal

HXB2 Location gp160 (432–451)

Author Location gp120 (42-61 LAI)

Epitope KAMYAPPISGQIRCSSNITG

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia HIV component: Env

Species (Isotype) mouse

Ab Type gp120 C4

References Collado et al. 2000

• Vaccinia p14 can elicit NAbs and p39 tends to be immunodominant, so these two proteins were fused to regions of HIV-1 Env – reduced glycosylation was noted when p14 or p39 was placed in the N-term region of the fusion protein – chimeric proteins shifted the Env Ab response from V3 to either a C1 or C4 domain, depending on the construct – all chimeric Env proteins: 14kEnv, 39kEnv, and Env39k elicited a strong Ab response to the C1 region of gp120 (LFCASDAKAYDTEVHNVWAT), and Env39k mounted a strong response to the C4 region (KAMYAPPISGQIRCSSNITG) Collado et al. [2000]

No. 598

**MAb ID** 1662

**HXB2 Location** gp160 (433–439)

Author Location gp120 (IIIB)

Epitope AMYAPPI

Neutralizing no

Immunogen vaccine

Vector/Type: poliovirus HIV component:

Species (Isotype)

Ab Type gp120 C4

References McKeating et al. 1992b

 1662: Did not bind to native gp120, epitope not exposed. McKeating et al. [1992b]

No. 599

**MAb ID** 1663

HXB2 Location gp160 (433-439)

Author Location gp120 (IIIB)

Epitope AMYAPPI

Neutralizing no

Immunogen vaccine

Vector/Type: poliovirus HIV component:

Env

Species (Isotype) Species (Isotype) Ab Type gp120 C4 Ab Type gp120 C4 References McKeating et al. 1992b References McKeating et al. 1992b • 1663: Did not bind to native gp120, epitope not exposed. McK- • 1804: Did not bind to native gp120, epitope not exposed. McKeating et al. [1992b] eating et al. [1992b] No. 600 No. 604 **MAb ID** 1664 **MAb ID** 1807 HXB2 Location gp160 (433-439) HXB2 Location gp160 (433-442) Author Location gp120 (IIIB) Author Location gp120 (IIIB) Epitope AMYAPPISGQ Epitope AMYAPPI Neutralizing no Neutralizing no Immunogen vaccine Immunogen vaccine Vector/Type: poliovirus HIV component: Vector/Type: poliovirus HIV component: Env Species (Isotype) Species (Isotype) Ab Type gp120 C4 Ab Type gp120 C4 References McKeating et al. 1992b References McKeating et al. 1992b • 1664: Did not bind to native gp120, epitope not exposed. McK- • 1807: Did not bind to native gp120, epitope not exposed. McKeating *et al.* [1992b] eating et al. [1992b] No. 601 No. 605 **MAb ID** 1697 **MAb ID** 1808 HXB2 Location gp160 (433–439) **HXB2 Location** gp160 (433–442) Author Location gp120 (IIIB) Author Location gp120 (IIIB) Epitope AMYAPPI Epitope AMYAPPISGO Neutralizing no Neutralizing no Immunogen vaccine Immunogen vaccine Vector/Type: poliovirus HIV component: Vector/Type: poliovirus HIV component: Species (Isotype) Species (Isotype) Ab Type gp120 C4 Ab Type gp120 C4 References McKeating et al. 1992b References McKeating et al. 1992b • 1697: Did not bind to native gp120, epitope not exposed. McK- • 1808: Did not bind to native gp120, epitope not exposed. McKeating *et al*. [1992b] eating et al. [1992b] No. 602 No. 606 **MAb ID** 1794 MAb ID polyclonal (VEI5) HXB2 Location gp160 (433-442) HXB2 Location gp160 (454–474) Author Location gp120 (IIIB) **Author Location** Env Epitope AMYAPPISGQ Epitope LTRDGGNNNNESEIFRPGGGD Neutralizing no **Neutralizing** Immunogen vaccine Immunogen HIV-1 infection Vector/Type: poliovirus HIV component: Species (Isotype) human **Ab Type** gp120 V1, gp120 V2, gp120 V3, gp120 V4, Species (Isotype) gp120 V5 Ab Type gp120 C4 References Carlos et al. 1999 References McKeating et al. 1992b • Antibody response to the epitopes in a vaccine construct (VEI) • 1794: Did not bind to native gp120, epitope not exposed. McKcontaining peptides from 5 hypervariable regions of gp120 was eating et al. [1992b] detected in the sera of HIV-1 positive subjects, including sera from 6 non-subtype B infections – serum samples from San **No.** 603 Francisco, Canada and Puerto Rico cohort showed presence of **MAb ID** 1804 antibodies against all five VEI hypervariable regions, but most HXB2 Location gp160 (433–442) consistently against the V3 region peptide NNNTRKSIRIGP-Author Location gp120 (IIIB) GRAFYTTGDIGNIRQ. Carlos et al. [1999] Epitope AMYAPPISGQ **No.** 607 Neutralizing no MAb ID polyclonal Immunogen vaccine *Vector/Type:* poliovirus *HIV component:* **HXB2 Location** gp160 (460–467) Author Location gp120 (LAI)

**Epitope NNNNGSEI** 

Subtype B

Neutralizing

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

**Species (Isotype)** human **Ab Type** gp120 V5

**References** Loomis-Price *et al.* 1997

• HIV-1 + positive individuals were given a gp160 vaccine as immunotherapy, and this region was the most reactive new epitope as measured by a modified Pepscan technique which improved sensitivity – 4/14 showed vaccine-induced reactivity. Loomis-Price *et al.* [1997]

No. 608

MAb ID CRA1 (CRA-1, CRA1(ARP 323))

**HXB2 Location** gp160 (461–470)

Author Location gp120 (451–470 LAI)

**Epitope** SNNESEIFRL

Subtype B Neutralizing no Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 V5-C5

Research Contact M. Page, NIBSC, UK

**References** Koefoed *et al.* 2005; Yang *et al.* 2000; Trkola *et al.* 1996a; Moore & Sodroski 1996; Moore *et al.* 1994c; Moore *et al.* 1994d; Moore & Ho 1993

**Keywords** antibody binding site definition and exposure

- CRA1: UK Medical Research Council AIDS reagent: ARP323.
- CRA1: IgG antibody phage display libraries were created from HIV-1 + individuals after pre-selection of PBMC with gp120, as an alternative to using bone marrow for generating libraries. CRA1 was among a set of Abs used for competition studies to define the binding sites of the newly isolated MAbs, representing a MAb with a linear C5 epitope. Koefoed *et al.* [2005] (antibody binding site definition and exposure)
- CRA1: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang *et al.* [2000]
- CRA1: C5 region linear epitope, binds weakly to nondenatured monomeric gp120 – reciprocal binding inhibition with anti-C5 antibodies 1C1 and M91 – non-reciprocal binding enhancement some C1 and V2 antibodies – non-reciprocal binding inhibition of some CD4 binding site antibodies. Moore & Sodroski [1996]
- CRA1: Does not neutralize JR-FL nor block gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]

 CRA1: Some C5 mutations abrogate binding 470 P/L or G, 475 M/S, some C2 mutations enhance binding. Moore et al. [1994d]

- CRA1: The relative affinity for denatured/native gp120 is 24

   C5 mutations 470 P/L or G, 475 M/S impairs binding to the native gp120 only mutation 470 P/L impairs binding to denatured. Moore *et al.* [1994c]
- CRA1: Bound preferentially to denatured IIIB and SF2 gp120.
   Moore & Ho [1993]

**No.** 609

MAb ID M91

**HXB2 Location** gp160 (461–470) **Author Location** gp120 (451–470 LAI)

**Epitope SNNESEIFRL** 

Subtype B Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) rat (IgG2a)

Ab Type gp120 V5-C5

Research Contact Fulvia di Marzo Veronese

**References** Zwick *et al.* 2003; Yang *et al.* 2000; Binley *et al.* 1998; Ditzel *et al.* 1997; Moore & Sodroski 1996; Moore *et al.* 1994d; Moore *et al.* 1994c; di Marzo Veronese *et al.* 1992

Keywords antibody interactions

- M91: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a C5-binding MAb that had no impact on 4KG5 binding. Zwick *et al.* [2003] (antibody interactions)
- M91: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang *et al.* [2000]
- M91: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer. Binley et al. [1998]
- M91: C5 region linear epitope, binds weakly to nondenatured monomeric gp120 – M91 binding was enhanced by 1C1, but 1C1 binding was inhibited by M91 – non-reciprocal binding enhancement of C1 and V2 antibodies – non-reciprocal binding inhibition of CD4 binding site antibodies. Moore & Sodroski [1996]

- mutation in position 470 P/L impairs binding. Moore et al. [1994c]

- M91: 470 P/L impairs binding, but not 475 D/V, in contrast to CRA1 – some C2 mutations can enhance binding. Moore et al. [1994d]
- M91: Immunoblot reactive, RIP negative, but precipitates deglycosylated gp120 - reacts with strains IIIB, 451, MN, RF, and RUTZ. di Marzo Veronese et al. [1992]

**No.** 610 **MAb ID** 9201

HXB2 Location gp160 (471–482)

Author Location gp120 (475–486 LAI)

Epitope GGGDMRDNWRSE

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: peptide

Species (Isotype) mouse (IgG1)

Ab Type gp120 C5

Research Contact Du Pont de Nemours, Boston, MA

References Dairou et al. 2004; McDougal et al. 1996

Keywords antibody binding site definition and exposure

- 9201: This paper describes a slightly different epitope, stating 9201 was raised against the peptide MRDNWRSELIKY, located within the alpha 5 helix in the C5 terminal region of gp120. Two MAbs were used to determine the photodamage location in HIV-1 Env induced by sulfonated anionic porphyrins. The negatively charged porphyrins interact with positive charge in the V3 loop. When light activated, they damage amino acid side chains in the C5 region of Env, as evidenced by inhibition of binding of C5 MAb 9201, but not V3 MAb 13105100. Anionic porphyrins could be used in targeted photodynamic decontamination of biological fluids, such as blood, killing HIV Research Contact S. Nigida, NCI, USA without disabling the function of desirable transfusion products. Dairou et al. [2004] (antibody binding site definition and exposure)
- 9201: Does not neutralize LAI. This paper notes the peptide binding region is GGGDMRDNWRSE. McDougal et al. [1996] (antibody binding site definition and exposure)

No. 611

MAb ID 1C1

HXB2 Location gp160 (471–490)

Author Location gp120 (471–490 LAI)

Epitope GGGDMRDNWRSELYKYKVVK

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI Research Contact S. Ranjbar, NIBSC, UK

HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C5

Research Contact Repligen Inc, Cambridge, MA, commercial

References Zwick et al. 2003; Moore & Sodroski 1996;

VanCott et al. 1995; Moore et al. 1994d;

Moore et al. 1994c

**Keywords** antibody interactions

- M91: The relative affinity for denatured/native gp120 is 24 1C1: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a C5-binding MAb that had no impact on 4KG5 binding. Zwick et al. [2003] (antibody interactions)
  - 1C1: C5 region linear epitope, binds weakly to nondenatured monomeric gp120 - M91 binding was enhanced by 1C1, but 1C1 binding was inhibited by M91 - non-reciprocal binding enhancement of some C1 and V2 antibodies - non-reciprocal binding inhibition of some CD4 binding site antibodies. Moore & Sodroski [1996]
  - 1C1: Linear epitope not exposed on conformationally intact gp120. VanCott et al. [1995]
  - 1C1: The relative affinity for denatured/native gp120 is 15. Moore *et al.* [1994c]
  - 1C1: C2 and V3 regions substitutions can influence binding. Moore et al. [1994d]

No. 612

MAb ID 3F5

**HXB2 Location** gp160 (471–490)

Author Location gp120 (471–490 LAI)

Epitope GGGDMRDNWRSELYKYKVVK

Subtype B

Neutralizing

Immunogen vaccine

Strain: B clade LAI HIV component: Env

Species (Isotype) mouse (IgG)

Ab Type gp120 C5

References Moore et al. 1994c

• 3F5: The relative affinity for denatured/native gp120 is 100. Moore et al. [1994c]

No. 613

**MAb ID** 5F4/1

**HXB2 Location** gp160 (471–490)

Author Location gp120 (471-490 LAI)

Epitope GGGDMRDNWRSELYKYKVVK

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: peptide Strain: HIV-2 ROD

Species (Isotype) mouse

Ab Type gp120 C5

References Moore et al. 1994c

• 5F4/1: V5-C5 region - preferentially binds SDS-DTT denatured gp120 (>10 fold) - mutation 485 K/V impairs binding. Moore et al. [1994c]

No. 614

**MAb ID** 660-178

HXB2 Location gp160 (471–490) Author Location gp120 (471–490 LAI)

**Epitope** GGGDMRDNWRSELYKYKVVK • B221: Called 221 – C2 and V3 substitutions influence binding. Subtype B Moore et al. [1994d] **Neutralizing** • B221: Called 221 - bound preferentially to denatured IIIB Immunogen vaccine gp120. Moore & Ho [1993] Vector/Type: protein Strain: B clade LAI No. 617 HIV component: Env MAb ID 8C6/1 Species (Isotype) mouse (IgG) HXB2 Location gp160 (471–490) Ab Type gp120 C5 Author Location gp120 (471-490 LAI) Research Contact G. Robey, Abbott Labs Epitope GGGDMRDNWRSELYKYKVVK References Moore et al. 1994d; Moore et al. 1994c Subtype B • 660-178: The relative affinity for denatured/native gp120 is **Neutralizing** >100. Moore et al. [1994c] Immunogen vaccine • 660-178: DeltaV1/V2 and DeltaV1/V2/V3 reduce binding -Strain: B clade LAI C2 and C5 mutations enhance binding. Moore et al. [1994d] Species (Isotype) mouse (IgG) Ab Type gp120 V5-C5 No. 615 **MAb ID** 9301 Research Contact S. Ranjbar, NIBSC, UK HXB2 Location gp160 (471-490) References Moore et al. 1994c Author Location gp120 (471–490 LAI) • 8C6/1: UK Medical Research Council AIDS reagent: Epitope GGGDMRDNWRSELYKYKVVK ARP3052. Subtype B • 8C6/1: V5-C5 region - preferentially binds SDS-DTT dena-Neutralizing tured gp120 (>30 fold) - mutation 485 K/V impairs binding. Moore et al. [1994c] Immunogen vaccine Vector/Type: protein Strain: B clade LAI No. 618 HIV component: Env MAb ID H11 Species (Isotype) mouse (IgG) HXB2 Location gp160 (472–477) Ab Type gp120 C5 Author Location gp120 (472–477 HXB2) Research Contact Dupont, commercial Epitope GGDMRD References Wagner et al. 1996; Moore et al. 1994d; Subtype B Moore et al. 1994c; Moore & Ho 1993; Skin-**Neutralizing** ner et al. 1988b Immunogen • 9301: Wagner et al. claim that Nea 9301 is anti-V3 - might Species (Isotype) mouse they have meant MAb 9305? Wagner et al. [1996] Ab Type gp120 C5 • 9301: The relative affinity for denatured/native gp120 is 19. References Pincus et al. 1996; Pincus & McClure 1993 Moore et al. [1994d] • H11: Binds to gp120 but not to infected cells – when linked to • 9301: Bound preferentially to denatured IIIB gp120. Moore & ricin A, the immunotoxin did not mediate cell killing – sCD4 Ho [1993] has no effect. Pincus & McClure [1993]; Pincus et al. [1996] **No.** 616 No. 619 **MAb ID** B221 (221) MAb ID W2 HXB2 Location gp160 (471–490) HXB2 Location gp160 (472–491) **Author Location** gp120 (471–490 LAI) Author Location gp120 (472-491 LAI) Epitope GGGDMRDNWRSELYKYKVVK Epitope GGDMRDNWRSELYKYKVVKI Subtype B Subtype B **Neutralizing** Neutralizing Immunogen vaccine Immunogen vaccine Vector/Type: protein Strain: B clade NL43 Strain: B clade LAI HIV component: Env HIV component: gp160 Species (Isotype) mouse (IgG) **Species** (**Isotype**) mouse ( $IgG1\kappa$ ) Ab Type gp120 C5 Ab Type gp120 C5 Research Contact D. Weiner, U. Penn., USA Research Contact Rod Daniels References Moore et al. 1994c References Moore et al. 1994d; Moore et al. 1994c; Bris-• W2: The relative affinity for denatured/native gp120 is 30 tow et al. 1994; Moore & Ho 1993 mutation 485 K/V impairs binding. Moore et al. [1994c] • B221: UK Medical Research Council AIDS reagent: ARP301. • B221: MAb generated in a study of the humoral immune re-No. 620 sponse to Baculovirus-expressed mis-folded rgp160 IIIB:NL43, MAb ID M38 MicroGenSys. Bristow et al. [1994] HXB2 Location gp160 (485–504) • B221: The relative affinity for denatured/native gp120 is 12 –

mutation 477 D/V impairs binding. Moore et al. [1994c]

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Epitope KYKVVKEIPLGVAPTKAKRR

**Author Location** gp120 (490–508)

Neutralizing no Immunogen vaccine

Vector/Type: virus Strain: B clade IIIB

HIV component: HIV-1

Species (Isotype) mouse

Ab Type gp120 C5

**References** Maksiutov *et al.* 2002; Beretta & Dalgleish 1994; DeSantis *et al.* 1994; Lopalco *et al.* 1993; Grassi *et al.* 1991; Beretta *et al.* 1987

- M38: This epitope is similar to a fragment of the human protein mast/stem cell growth factor receptor precursor, VVPTKAD-KRRSV, as well as to a fragment of IFN-related IFRD2 (PC4-B) protein, ARTKARSRVRDKRA. Maksiutov et al. [2002]
- M38: Infected individuals have HLA class I-gp120 crossreactive antibodies. DeSantis et al. [1994]
- M38: Binds to the carboxy terminus of gp120, in a gp41 binding region, and also to denatured human HLAs (antigenic homology) Lopalco *et al.* [1993]
- M38: Binds to gp120 and to a 80 kd human protein expressed on a small fraction of mononuclear cells in the lymph nodes. Beretta *et al.* [1987]

No. 621

MAb ID Chim 1 (C-1)

**HXB2 Location** gp160 (487–493)

Author Location gp120 (492–498 HXB2)

Epitope KVVKEIP

Subtype B

Neutralizing

Immunogen

Species (Isotype) humanized chimpanzee

References Pincus et al. 1996; Pincus & McClure 1993

• Chim 1: Binds to gp120 but not to infected cells – when linked to ricin A, the immunotoxin did not mediate cell killing – sCD4 has no effect. Pincus & McClure [1993]; Pincus et al. [1996]

No. 622

MAb ID polyclonal

**HXB2 Location** gp160 (490–511)

Author Location gp120 (495–516 BRU)

Epitope KIEPLGVAPTKAKRRVVQREKR

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human

References Maksiutov et al. 2002; Hernandez et al. 2000

- This epitope is similar to a fragment of the human protein mast/stem cell growth factor receptor precursor, VVPTKAD-KRRSV, as well as to a fragment of IFN-related IFRD2 (PC4-B) protein, ARTKARSRVRDKRA. Maksiutov et al. [2002]
- Chimeric peptide combining two peptides gp160(495-516 and 584-612) served as a specific and broadly reactive antigen for diagnostic detection of HIV-1. Hernandez *et al.* [2000]

**No.** 623

**MAb ID** 110.1

HXB2 Location gp160 (491–500)

**Author Location** gp120 (491–500 LAI)

**Epitope IEPLGVAPTK** 

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: HIV infected-cell lysate Strain: B clade BRU HIV component:

HIV-1

**Species (Isotype)** mouse (IgG1 $\kappa$ )

**Ab Type** gp120 C5

References Maksiutov et al. 2002; Beretta & Dalgleish Research Contact Genetic Systems Corp, Seattle WA, E.

Kinney-Thomas

References Maksiutov et al. 2002; Valenzuela et al. 1998;

Binley *et al.* 1997a; McDougal *et al.* 1996; Cook *et al.* 1994; Moore *et al.* 1994c; Callahan *et al.* 1991; Pincus *et al.* 1991; Thomas *et al.* 1988; Linsley *et al.* 1988; Gosting *et al.* 

1987

**Keywords** antibody binding site definition and exposure, immunotoxin

- 110.1: There is another antibody with this ID that binds to gp120, but at aa 200-217.
- 110.1: This epitope is similar to a fragment of the human protein mast/stem cell growth factor receptor precursor, VVP-TKADKRRSV. Maksiutov et al. [2002]
- 110.1: Does not effect LAI viral binding or entry into CEM cells. Valenzuela *et al.* [1998]
- 110.1: Does not neutralize HIV-1 LAI. McDougal et al. [1996]
- 110.1: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon MAbs against the carboxy-terminus of gp120 inhibit gp120 binding to GalCer but not as potently as anti-V3 MAbs binding of GalCer to gp120 does not inhibit MAb binding. Cook *et al.* [1994]
- 110.1: The relative affinity for denatured/native gp120 is 0.7. Moore *et al.* [1994c] (antibody binding site definition and exposure)
- 110.1: Polyanionic polysaccharides were proposed to inhibit viral functions such as binding and syncytia formation through interations mediated through the local high positive charge density in the V3 loop. The binding of this antibody is not inhibited by dextransulfate, in contrast to anti-V3 antibodies. Callahan *et al.* [1991]
- 110.1: Difference was noted in the epitope: mapped to aa 421-429 (KQIINMWQE), the T1 sequence poor efficacy as an immunotoxin when linked to RAC. Pincus et al. [1991] (antibody binding site definition and exposure, immunotoxin)
- 110.1: Referred to as 110-1 does not inhibit CD4-gp120 binding or neutralize HIV-1 strains. Linsley *et al.* [1988]

**No.** 624

MAb ID 42F

**HXB2 Location** gp160 (491–500)

Author Location gp120 (491-500 HXB2)

Epitope IEPLGVAPTK

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 C5

References Maksiutov et al. 2002; Alsmadi & Tilley

1998; Alsmadi et al. 1997

 42F: This epitope is similar to a fragment of the human protein mast/stem cell growth factor receptor precursor, VVPTKAD-KRRSV. Maksiutov et al. [2002]

- 42F: A study of 6 anti-Env MAbs and their ability to bind or direct ADCC against target cells infected with IIIB, MN, SF-2, and RF – bound and directed lysis against strains IIIB, MN, SF-2, and RF, but not a clone of MN. Alsmadi & Tilley [1998]
- 42F: 42F and 43F were isolated from a long term non-progressor by EBV transformation of PBMC samples were taken 14 months apart both MAbs stained diverse strains of infected cells and directed ADCC were more potent for ADCC if the cell was infected with HIV-1, rather than just presenting absorbed gp120. Alsmadi et al. [1997]

No. 625

MAb ID 43F

HXB2 Location gp160 (491-500)

Author Location gp120 (491–500 HXB2)

**Epitope** IEPLGVAPTK

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\lambda$ )

Ab Type gp120 C5

References Maksiutov et al. 2002; Alsmadi et al. 1997

- 43F: This epitope is similar to a fragment of the human protein mast/stem cell growth factor receptor precursor, VVPTKAD-KRRSV. Maksiutov et al. [2002]
- 43F: 42F and 43F were isolated from a long term non-progressor by EBV transformation of PBMC samples were taken 14 months apart both MAbs stained diverse strains of infected cells and directed ADCC were more potent for ADCC if the cell was infected with HIV-1, rather than just presenting absorbed gp120. Alsmadi et al. [1997]

No. 626

**MAb ID** RV110026

HXB2 Location gp160 (491–500)

Author Location gp120 (491–500 LAI)

**Epitope** IEPLGVAPTK

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: peptide Strain: B clade LAI

Species (Isotype) human

Ab Type gp120 C5

Research Contact Commercial, Olympus Inc

**References** Maksiutov *et al.* 2002; Moore *et al.* 1994d; Moore *et al.* 1994c

- RV110026: This epitope is similar to a fragment of the human protein mast/stem cell growth factor receptor precursor, VVPTKADKRRSV. Maksiutov et al. [2002]
- RV110026: Preferentially binds SDS-DTT denatured gp120 (15 fold using R1/87 as capture reagent) Moore *et al.* [1994c]

No. 627

**MAb ID** 105-306

**HXB2 Location** gp160 (492–500)

Author Location gp120 (498–505 HAM112, O group)

Epitope KPFSVAPTP

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: O group

HAM112 HIV component: gp160

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type C-term

References Scheffel et al. 1999

• 105-306: Overlapping peptides based on group O HAM112 Env were tested for MAb reactivity – 105-306 bound to two overlapping peptides. Scheffel *et al.* [1999]

No. 628

MAb ID GV1G2

HXB2 Location gp160 (494-499)

Author Location gp120 (494-499 IIIB)

Epitope LGVAPT

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein-Ab complex HIV com-

ponent: gp120-Mab complex

Species (Isotype) mouse

Ab Type gp120 C5

References Denisova et al. 1996

GV1G2: When anti-V3 MAb M77 was bound to gp120 and used as an immunogen, it stimulated many MAbs to linear epitopes – MAbs GV12F6 and GV3H1 are homologous to GV1G2 and were generated in the same experiment. Denisova et al. [1996]

No. 629

**MAb ID** 750-D

**HXB2 Location** gp160 (498–504)

**Author Location** gp120 (503–509)

**Epitope** PTKAKRR

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG $3\lambda$ )

Ab Type C-term

References Hioe et al. 2000; Forthal et al. 1995

- 750-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses
   CD4BS MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells C5 MAbs 450-D and 750-D did not effect proliferation. Hioe et al. [2000]
- 750-D: Not neutralizing, positive ADCC activity, and no viral enhancing activity. Forthal et al. [1995]

**No.** 630

**MAb ID** 450-D (450-D-3, 450D)

HXB2 Location gp160 (498–504)

**Author Location** gp120 (475–486 BH10)

Epitope PTKAKRR(orRRVVQRE,orMRDNWRSELYKY-

dependingonreference)

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human ( $IgG1\lambda$ )

Ab Type gp120 C5

Research Contact Susan Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu), NYU Med

Center, NY, NY

References Verrier et al. 2001; Hioe et al. 2001; Hioe et al. 2000; Hioe et al. 1997b; Li et al. 1997; Manca et al. 1995a; Forthal et al. 1995; Cook et al. 1994; Gorny et al. 1994; Laal et al. 1994; Spear et al. 1993; Karwowska et al. 1992b; Karwowska et al. 1992a; Durda et al. 1988

- 450-D: CD4BS MAbs when complexed with gp120, inhibit proliferation of gp120-specific CD4 T-cells and IFN gamma Research Contact Susan production – 450-D does not have this effect and was used as a control in this study. Hioe et al. [2001]
- 450-D: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 – six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 – no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001]
- 450-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses - CD4BS MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells - C5 MAbs 450-D and 750-D did not effect proliferation. Hioe et al. [2000]
- 450-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs - BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) - isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations – US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe et al. [1997b]
- 450-D: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env - 50% neutralization could not be achieved at a maximal concentration of 6 mug/ml. Li et al. [1997]
- 450-D: No neutralizing activity, no ADCC activity, and no viral enhancing activity. Forthal et al. [1995]
- 450-D: Virions complexed to gp120 Ab facilitate presentation of p66 RT epitopes to Th cells. Manca et al. [1995a]
- 450-D: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon – MAbs against the carboxy-terminus of gp120 do not inhibit gp120 binding to GalCer - binding of GalCer to gp120 does not inhibit MAb binding. Cook et al. [1994]
- 450-D: Epitope is defined as PTKAKRR. Gorny et al. [1994]
- 450-D: Not neutralizing alone, could synergize anti-CD4 binding site antibody neutralization. Laal et al. [1994]
- 450-D: Did not mediate deposition of complement component C3 on HIV infected cells. Spear et al. [1993]
- 450-D: Bound to MN, SF-2 and IIIB, but was not neutralizing. Karwowska et al. [1992a]

No. 631 **MAb ID** 670-D (670)

HXB2 Location gp160 (498–504) Author Location gp120 (503-509) **Epitope** PTKAKRR

Neutralizing no

**Immunogen** HIV-1 infection **Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 C5

Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu), NYU, NY

References Gorny et al. 2005; Zwick et al. 2003; Verrier et al. 2001; Nyambi et al. 2000; Gorny & Zolla-Pazner 2000; Altmeyer et al. 1999; Nyambi et al. 1998; Gorny et al. 1998; Hioe et al. 1997b; Gorny et al. 1997; Hill et al. 1997; Forthal et al. 1995; Zolla-Pazner et al. 1995

## **Keywords** antibody interactions

- 670: 2909 is a human anti-Env NAb that was selected by neutralization assay and binds to the quaternary structure on the intact virion. ELISA-based competition assays and subsequent mutational analysis determined that the CD4BS and V2 and V3 loops contribute to the 2909 epitope: 2909 binding was inhibited by MAbs 447-52d (anti-V3), 830A (anti-V2), and IgG1b12 (anti-CD4BS) and sCD4. 2909 was not inhibited by MAbs 670, 1418, nor 2G12. Gorny et al. [2005]
- 670-D: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a C5-binding MAb that had no impact on 4KG5 binding. Zwick et al. [2003] (antibody interactions)
- 670-D: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 – no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001]
- 670-D: A gp120 C5 MAb used as a negative control in a study of anti-gp41 MAbs. Gorny & Zolla-Pazner [2000]
- 670-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 4 C5 MAbs, 2 bound well, 2 bound weakly - MAb 670-D bound 21/26, and was the most cross-reactive C5 MAb. Nyambi et al. [2000]
- 670-D: A Semliki Forest virus (SFV) expression system carrying BX08 env was used to study the conformation of gp120 intracytoplasmic gp120 was recognized by the anti-V3 MAbs K24 and F5.5, while gp120 at the plasma membrane was detected only by conformation dependent MAbs 2G12, 670-D and 694/98D and not V3 MAbs - expression in rat brain also showed that surface-expressed Env was recognized only by the conformation-dependent antibodies and not by anti-V3 antibodies. Altmeyer et al. [1999]

- 670-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H anti-C5 Abs 670-D and 1331A bound to 3/4 B clade viruses (they didn't bind to IIIB), and to subtype D MAL 670-D also reacted with subtype A. Nyambi *et al.* [1998]
- 670-D: gp120 can inhibit MIP-1alpha from binding to CCR5, but this inhibitory effect is blocked by pre-incubation of gp120 with three anti-V3 MAbs: 447, 257, 1027 MAb 670 which binds in the C5 region had no effect. Hill *et al.* [1997]
- 670-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe *et al.* [1997b]
- 670-D: Not neutralizing, positive ADCC activity, and no viral enhancing activity, numbering provided suggests epitope is RRVVQRE. Forthal et al. [1995]
- 670-D: Group specific cross-clade binding in serotyping study using flow-cytometry. Zolla-Pazner et al. [1995]

**No.** 632

**MAb ID** 158F3

HXB2 Location gp160 (499-511)

Author Location gp120 (BaL)

Epitope TKAKRRVVQREKR

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: gp120-CD4 complex HIV component: gp120 Adjuvant: Ribi adjuvant

(MPL+TDM) (RIBI)

**Species** (**Isotype**) humanized mouse ( $IgG2\kappa$ )

Ab Type C-term

Research Contact Abraham Pinter, Lab. of Retrovirology, Public Research Institute, pinter@phri.org

References He et al. 2003

**Keywords** antibody binding site definition and exposure, vaccine antigen design

• 158F3: Vaccination of a gp120-CD4 fusion complex in six transgenic XMG2 XenoMouse mice that produce human IgG2 with K light chain did not produce any neutralizing antibodies. 36/39 MAbs derived from one of these mice were in one of two competition groups that were conformational and specific for the complex, suggesting this chimeric vaccine may be of little value, as immunodominant responses are to epitopes unique to the complex, not present in native Env. This MAb is one of the 3/36 non-neutralizing MAbs that bound to linear epitopes on gp120. He *et al.* [2003] (antibody binding site definition and exposure, vaccine antigen design)

No. 633

**MAb ID** 161D7

**HXB2 Location** gp160 (499–511)

Author Location gp120 (BaL)

**Epitope** TKAKRRVVQREKR

Subtype B Neutralizing no

Immunogen vaccine

Vector/Type: gp120-CD4 complex HIV

component: gp120 Adjuvant: Ribi adjuvant

(MPL+TDM) (RIBI)

**Species (Isotype)** humanized mouse (IgG2 $\kappa$ )

Ab Type C-term

- BZ167 was the only isolate inhibited by all polyclonal sera Research Contact Abraham Pinter, Lab. of Retrovirology, Pub-

lic Research Institute, pinter@phri.org

References He et al. 2003

**Keywords** antibody binding site definition and exposure,

vaccine antigen design

• 161D7: Vaccination of a gp120-CD4 fusion complex in six transgenic XMG2 XenoMouse mice that produce human IgG2 with K light chain did not produce any neutralizing antibodies. 36/39 MAbs derived from one of these mice were in one of two competition groups that were conformational and specific for the complex, suggesting this chimeric vaccine may be of little value, as immunodominant responses are to epitopes unique to the complex, not present in native Env. This MAb is one of the 3/36 non-neutralizing MAbs that bound to linear epitopes on gp120. He *et al.* [2003] (antibody binding site definition and exposure, vaccine antigen design)

**No.** 634

MAb ID polyclonal

**HXB2 Location** gp160 (503–509)

**Author Location** gp120 (471–477)

Epitope RRVVQRE

Neutralizing

Immunogen vaccine

Vactor/Type pontide

Vector/Type: peptide HIV component:

gp120

Species (Isotype) mouse (IgG)

References Jeyarajah et al. 1998

Mice were immunized with peptide APTKAKRRVVQREKR –
epitope excision and extraction combined with mass spectrometry was used to map the fine structure of epitopes recognized by polyclonal Ab to HIV-1 Env – a major epitope was identified between positions 472 and 478. Jeyarajah et al. [1998]

No. 635

**MAb ID** 722-D

HXB2 Location gp160 (503–509)

**Author Location** gp120 (503–509)

Epitope RRVVQRE

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type C-term

References Forthal et al. 1995; Laal et al. 1994

- 722-D: No neutralizing activity, no ADCC activity, and no viral enhancing activity. Forthal et al. [1995]
- 722-D: Not neutralizing alone, could synergize anti-CD4 binding site antibody neutralization. Laal et al. [1994]

**No.** 636

MAb ID polyclonal

HXB2 Location gp160 (503-511)

Author Location gp120 (508-516)

Epitope RRVVQREKR

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type C-term

References Loomis-Price et al. 1997; Palker et al. 1987

• Most HIV-1 + individuals have an antibody response to this epitope – in this study, reactivity to RRVVQREKR was used as a positive control for HIV-1 + gp160 vaccine recipients. Loomis-Price *et al.* [1997]

No. 637

**MAb ID** 1331A

**HXB2 Location** gp160 (503–511)

Author Location gp120 (510-516)

Epitope dwVVQREKR

**Neutralizing** 

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG $3\lambda$ )

Ab Type gp120 C5

Research Contact Susan

Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu) (NYU Med.

Center)

References Zwick et al. 2003; Edwards et al. 2002; Gorny et al. 2002; Nyambi et al. 2000; Hochleitner et al. 2000b; Gorny et al. 2000; Nyambi et al. 1998

- 1331A: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a C5-binding MAb that had no impact on 4KG5 binding. Zwick et al. [2003]
- 1331A: Truncation of the gp41 cytoplasmic domain of X4, R5, and X4R5 viruses forces a conformation that more closely resembles the CD4 bound state of the external Envelope, enhancing binding of CD4i MAbs 17b and 48d and of CD4BS MAbs F105, b12, and in most cases of glycosylation site dependent MAb 2G12 and the anti-gp41 MAb 246D in contrast, binding of the anti-V2 MAb 697D and the anti-V3 MAb 694/98D were not affected viruses bearing the truncation were more sensitive to neutralization by MAbs 48d, b12, and 2G12 the anti-C5 MAb 1331A was used to track levels of cell surface expression of the mutated proteins. Edwards *et al.* [2002]
- 1331A: Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated the six new MAbs all bind to the tip of the V3 loop and cross-compete with the MAb 447-52D and are conformationally sensitive MAbs showed cross-clade binding to native, intact virions and the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization

studies), 654 (anti-CD4BS used as a conformation-sensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control as binding was not diminished by treating gp120 with DTT or sodium metaperiodate to reduce disulfide bonds), and MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades). Gorny *et al.* [2002]

- 1331A: Core epitope dwVVQREKR maps to gp120(510-516)

   binding of panel of 21 MAbs to soluble oligomeric gp140
   versus gp41 or gp120 monomers was compared no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer C5 MAbs 858-D, 989-D and 1331A bound with a 5-10 fold preference for the monomer. Gorny et al. [2000]
- 1331A: The Ab binding site was studied with epitope excision (protein is bound in native conformation to immobilized MAb, then digested with proteolytic enzymes) and extraction (protein is digested then allowed to react with Ab), followed by mass spectroscopy two non-contiguous aa in C5 were protected, E-507 and I-487, which are thought to be located on opposite sides of hydrophobic pocket involved in gp120/gp41 interaction. Hochleitner *et al.* [2000b]
- 1331A: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 4 C5 MAbs, 2 bound well, 2 bound weakly – MAb 858-D bound only 4/26, the worst of all anti-C5 MAbs tested, while MAb 1331A, which shares the same core epitope (positions 495-516), bound to 18/26. Nyambi et al. [2000]
- 1331A: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H anti-C5 Abs 670-D and 1331A bound to 3/4 B clade viruses (they don't bind to IIIB), and to subtype D MAL. Nyambi *et al.* [1998]

**No.** 638

**MAb ID** 1131-A

**HXB2 Location** gp160 (505–511)

Author Location gp120 (510-516 LAI)

Epitope VVQREKR

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG $3\lambda$ )

Ab Type C-term

**References** Bandres *et al.* 1998

• 1131-A: A very high affinity antibody used in studies that demonstrate that CXCR4 can bind to gp120 in the absence of CD4-gp120 interactions, and that this binding can be enhanced by Env deglycosylation. Bandres *et al.* [1998]

**No.** 639

**MAb ID** 858-D

**HXB2 Location** gp160 (505–511)

Author Location gp120 (510-516 LAI)

Epitope VVQREKR

Subtype B

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

Ab Type C-term

Research Contact Susan Zolla-Pazner las01@mcrcr6.med.nyu) (NYU Med. Center)

> References Nyambi et al. 2000; Gorny et al. 2000; Forthal et al. 1995; Zolla-Pazner et al. 1995

- 858-D: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared – no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer - C5 MAbs 858-D, 989-D and 1331A bound with a 5-10 fold preference for the monomer. Gorny et al. [2000]
- 858-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 4 C5 MAbs, 2 bound well, 2 bound weakly - MAb 858-D bound only 4/26, the worst of all anti-C5 MAbs tested, while MAb 1331A, which shares the same core epitope (positions 495-516), bound to 18/26 isolates. Nyambi *et al.* [2000]
- enhancing activity. Forthal et al. [1995]
- 858-D: Group specific cross-clade binding in serotyping study using flow-cytometry. Zolla-Pazner et al. [1995]

**No.** 640

**MAb ID** 989-D

HXB2 Location gp160 (505-511)

Author Location gp120 (LAI)

Epitope VVQREKR

Subtype B

**Neutralizing** 

Immunogen HIV-1 infection Species (Isotype) human (IgG)

Ab Type C-term

Research Contact Susan

Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

References Nyambi et al. 2000; Gorny et al. 2000; Zolla-Pazner et al. 1995

- 989-D: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared - no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer - C5 MAbs 858-D, 989-D and 1331A bound with a 5-10 fold preference for the monomer. Gorny et al. [2000]
- 989-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 4 C5 MAbs, 2 bound well, 2 bound weakly – MAb 989-D bound to 6/26 isolates. Nyambi et al. [2000]
- 989-D: In serotyping study using flow-cytometry, showed B clade specificity, but only reacted with 7/11 B clade virus. Zolla-Pazner et al. [1995]

No. 641

MAb ID 1A1

HXB2 Location gp160 (525-543)

**Author Location** gp41 (526–543 BH10)

Epitope AAGSTMGAASMTLTVQARQ

Neutralizing no

Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ )

(Zol- Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna,

References Maksiutov et al. 2002; Buchacher et al. 1994

- 1A1: This epitope is similar to a fragment of the HLA class II histocompatibility antigen, GGSCMAALTVTLTV. Maksiutov et al. [2002]
- 1A1: Human MAb generated using EBV transformation of PBL from HIV-1 + volunteers. Buchacher et al. [1994]

No. 642

MAb ID 24G3

HXB2 Location gp160 (525-543)

**Author Location** gp41 (526–543 BH10)

Epitope AAGSTMGAASMTLTVQARQ

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

• 858-D: No neutralizing activity, no ADCC activity, and no viral Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna, Austria

> References Maksiutov et al. 2002; Buchacher et al. 1994; Buchacher et al. 1992

- 24G3: This epitope is similar to a fragment of the HLA class II histocompatibility antigen, GGSCMAALTVTLTV. Maksiutov et al. [2002]
- 24G3: Human MAb generated by electrofusion of PBL from HIV-1 + volunteers with CB-F7 cells. Buchacher et al. [1994]

No. 643

MAb ID 25C2 (IAM 41-25C2)

**HXB2 Location** gp160 (525–543)

**Author Location** gp41 (526–543 BH10)

Epitope AAGSTMGAASMTLTVQARQ

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna, Austria and Viral Testing Systems, Houston,

References Maksiutov et al. 2002; Sattentau et al. 1995; Buchacher et al. 1994: Buchacher et al. 1992

- 25C2: This epitope is similar to a fragment of the HLA class II histocompatibility antigen, GGSCMAALTVTLTV. Maksiutov et al. [2002]
- 25C2: Called IAM 41-25C2 Binding domain overlaps sites that are critical for gp120-gp41 association - binding is enhanced by sCD4 - binding region defined as: gp41(21-38 BH10). Sattentau et al. [1995]
- 25C2: Human MAb generated by electrofusion of PBL from HIV-1 + volunteers with CB-F7 cells - binds oligomeric and monomeric gp41, and gp160. Buchacher et al. [1994]

No. 644

MAb ID 5F3

HXB2 Location gp160 (525-543)

**Author Location** gp41 (526–543 BH10)

Epitope AAGSTMGAASMTLTVQARQ

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna,

References Maksiutov et al. 2002; Buchacher et al. 1994

- 5F3: This epitope is similar to a fragment of the HLA class II histocompatibility antigen, GGSCMAALTVTLTV. Maksiutov et al. [2002]
- 5F3: Human MAb generated by electrofusion of PBL from HIV-1 + volunteers with CB-F7 cells. Buchacher *et al.* [1994]

No. 645

**MAb ID**  $\alpha(566-586)$ 

HXB2 Location gp160 (561-581)

Author Location gp41 (566-586 BRU)

Epitope AQQHLLQLTVWGIKQLQARIL

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

References Poumbourios et al. 1992

No. 646

MAb ID PC5009

**HXB2 Location** gp160 (572–591)

**Author Location** gp41 (577–596 BRU)

Epitope GIKQLQARILAVERYLKDQQ

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component:

gp160

Species (Isotype) mouse

References Poumbourios et al. 1992

 PC5009: Recognized only monomeric gp41. Poumbourios et al. [1992]

No. 647

MAb ID polyclonal α577-596

HXB2 Location gp160 (572–591)

**Author Location** gp41 (577–596 BRU)

Epitope GIKQLQARILAVERYLKDQQ

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Poumbourios et al. 1992

• alpha(577-596): Affinity purified from HIV-1 + plasma – preferentially bind oligomer. Poumbourios *et al.* [1992]

No. 648

MAb ID polyclonal

HXB2 Location gp160 (576-592)

Author Location gp41 (583-599)

Epitope LQARILAVERYLKDQQL

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Klasse et al. 1993b

42 HIV-1 positive human sera were tested against wildtype peptide, and peptide with substitution 589 A to T: 11/42 reacted strongly with wildtype, weakly with A589T – 31 reacted weakly with parental, even more weakly with substituted. Klasse et al. [1993b]

**No.** 649

MAb ID

HXB2 Location gp160 (577-583)

Author Location gp41 (582-589)

**Epitope QARILAV** 

Subtype B

Neutralizing yes

**Immunogen** HIV-1 exposed seronegative

Species (Isotype) human (IgA)

Ab Type Leucine zipper motif

References Clerici et al. 2002a

Six sera from HIV-exposed uninfected individuals(EU), HIV-infected individuals and healthy controls were analyzed for IgA Abs – neutralizing activity was observed with total IgA from both EU and HIV+ – the EU IgA exclusively bound to a distinctive epitope within gp41, QARILAV, in the coiled coil pocket important for gp120-gp41 interactions – sera of QAFILAV-immunized BALB/c mice was neutralizing with the dose-dependent behavior similar to 2F5. Clerici et al. [2002a]

No. 650

MAb ID

**HXB2 Location** gp160 (577–583)

**Author Location** gp41 (582–589)

**Epitope QARILAV** 

Subtype B

Neutralizing yes

Immunogen vaccine

Vector/Type: peptide HIV component: gp41 Adjuvant: Keyhole Limpit Haemocyanin (KLH)

Species (Isotype) mouse (IgA)

**Ab Type** Leucine zipper motif

References Clerici et al. 2002a

Six sera from HIV-exposed uninfected individuals(EU), HIV-infected individuals and healthy controls were analyzed for IgA Abs – neutralizing activity was observed with total IgA from both EU and HIV+ – the EU IgA exclusively bound to a distinctive epitope within gp41, QARILAV – sera of QAFILAV-immunized BALB/c mice was neutralizing with the dose-dependent behavior similar to 2F5. Clerici et al. [2002a]

No. 651

MAb ID 1F11

**HXB2 Location** gp160 (578–612)

Author Location gp41 (579–613 BH10)

Epitope ARILAVERYLKDQQLLGIWGCSGKLICTTAVP-

WNA

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna, Austria

**References** Gorny & Zolla-Pazner 2004; Buchacher *et al.* 1994; Buchacher *et al.* 1992

Keywords antibody generation, review

• 1F11: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (**review**)

• 1F11: Generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994] (antibody generation)

No. 652

MAb ID 1H5

HXB2 Location gp160 (578-612)

Author Location gp41 (579–613 BH10)

Epitope ARILAVERYLKDQQLLGIWGCSGKLICTTAVP-

WNA

Neutralizing no

Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ )

References Gorny & Zolla-Pazner 2004; Buchacher et al.

1994; Buchacher et al. 1992

Keywords antibody generation, review

• 1H5: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna, Zolla-Pazner [2004] (review)

• 1H5: Generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994] (antibody generation)

No. 653

MAb ID 3D9

**HXB2 Location** gp160 (578–612)

**Author Location** gp41 (579–613 BH10)

Epitope ARILAVERYLKDQQLLGIWGCSGKLICTTAVP-

WNA

Neutralizing no

**Immunogen** HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna,

References Gorny & Zolla-Pazner 2004; Buchacher et al.

1994; Buchacher et al. 1992

**Keywords** antibody generation, review

• 3D9: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)

• 3D9: Generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994] (antibody generation)

No. 654

MAb ID 4B3

HXB2 Location gp160 (578-612)

**Author Location** gp41 (579–613 BH10)

Epitope ARILAVERYLKDQQLLGIWGCSGKLICTTAVP-

WNA

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna,

Austria

References Gorny & Zolla-Pazner 2004; Chen et al.

1994b; Buchacher et al. 1994; Buchacher et al.

1992

Keywords antibody generation, review

• 4B3: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)

• 4B3: Generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994] (antibody generation)

No. 655

MAb ID 4D4

HXB2 Location gp160 (578–612) **Author Location** gp41 (579–613 BH10)

Epitope ARILAVERYLKDQQLLGIWGCSGKLICTTAVP-

WNA

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Austria and Viral Testing Systems, Houston,

References Gorny & Zolla-Pazner 2004; Binley et al. 1999; Sattentau et al. 1995; Chen et al. 1994b;

Buchacher et al. 1994; Buchacher et al. 1992

Keywords antibody binding site definition and exposure, antibody generation, review, vaccine antigen

design

• 4D4: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)

• 4D4: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits - a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen – SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 - nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120gp41 complexes. Binley et al. [1999] (antibody binding site definition and exposure, vaccine antigen design)

4D4: Generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994] (antibody generation)

No. 656

MAb ID 4G2

**HXB2 Location** gp160 (578–612)

**Author Location** gp41 (579–613 BH10)

Epitope ARILAVERYLKDQQLLGIWGCSGKLICTTAVP-

Neutralizing no

Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ )

Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna,

Austria

References Gorny & Zolla-Pazner 2004; Buchacher et al. 1994; Buchacher et al. 1992

Keywords antibody generation, review

- 4G2: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- 4G2: Generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994] (antibody generation)

No. 657

MAb ID polyclonal

HXB2 Location gp160 (579-589) Author Location gp41 (586–596 IIIB)

Epitope RILAVERYLKD

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Adjuvant: BSA

Species (Isotype) rabbit, mouse

Ab Type C-domain

References Xiao et al. 2000b

• Strong epitope-specific neutralizing antibody responses were induced using the peptide C(RILAVERYLKD)\_2-BSA, but not full gp160. Xiao et al. [2000b]

No. 658

MAb ID polyclonal

HXB2 Location gp160 (579-589)

Author Location gp41 (586–596)

Epitope RILAVERYLKD

Neutralizing

Immunogen vaccine

ponent: gp160 Adjuvant: BSA

Species (Isotype) rabbit

Ab Type N-term

References Lu et al. 2000b; Lu et al. 2000c

• High titer response to ELDKWA and RILAVERYLKD was observed upon vaccination with multiple-epitope vaccine CG-GPGRAFY-G-ELDKWA-G-RILAVERYLKD conjugated to BSA, a weak response to GPGRAFY - immunization with CG-(ELDKWA-GPGRAFY)\_2-K was also tried, yielding a strong Ab response to both ELDKWA and GPGRAFY – gp160 vaccination yielded strong Ab response but not to any of the peptides studied here. Lu et al. [2000c,b]

No. 659

MAb ID

HXB2 Location gp160 (579-599)

Author Location gp41 (586–606)

Epitope RILAVERYLKDQQLLGIWGCS

Subtype B

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

References Wang et al. 1986

**Keywords** assay standardization/improvement

· Immunoabsorbant peptide antigen RIAVERYLKDQQLLGI-WGCS was used in a solid-phase enzyme immunoassay (EIA)to detect gp41-specific Abs in sera of virtually all HIV-1 infected individuals tested, with no false positives. This one 21 amino acid long peptide is recognized by sera from almost all AIDS patients, can be easily synthesized and employed for serological testing for HIV infection. Wang et al. [1986] (assay standardization/improvement)

No. 660

MAb ID polyclonal

HXB2 Location gp160 (579–599)

Author Location gp41 (583–604)

Epitope RILAVERYLKDQQLLGIWGCS

Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: desia-

lylated gp160

Species (Isotype) rabbit

References Benjouad et al. 1993

• MAbs raised against desialylated HIV-1 gp160 cross-react with HIV-2 gp140 due to immunodominant conserved epitope in gp41. Benjouad et al. [1993]

No. 661

MAb ID 2A2/26

HXB2 Location gp160 (579–601)

Author Location gp41 (584–606 BRU)

Epitope RILAVERYLKDQQLLGIWGCSGK

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: gp41

Species (Isotype) mouse (IgG)

References Poumbourios et al. 1995; Poumbourios et al.

- Vector/Type: protein, polyepitope HIV com- 2A2/26: Delta 550-561 (Delta LLRAIEAQQHLL), a region important for oligomer formation diminishes binding, Delta (550-561 +571-581) abrogates binding. Poumbourios et al. [1995]
  - 2A2/26: Immunodominant region, binds both oligomer and monomer. Poumbourios et al. [1992]

No. 662

**MAb ID** 50-69 (SZ-50.69, 50-69D)

HXB2 Location gp160 (579–603)

**Author Location** gp41 (579–603 BH10)

Epitope RILAVERYLKDQQLLGIWGCSGKLI

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG2\kappa$ )

Ab Type gp41 cluster I

Research Contact Susan Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu), NYU, NY

**References** McCaffrey *et al.* 2004; Ling *et al.* 2004; Gorny & Zolla-Pazner 2004; Finnegan *et al.* 

2002; Follis et al. 2002; Verrier et al. 2001; Zwick et al. 2001b; Nyambi et al. 2000; Gorny et al. 2000; Gorny & Zolla-Pazner 2000; Mitchell et al. 1998; Hioe et al. 1997b; Boots et al. 1997; Stamatatos et al. 1997; Klasse & Sattentau 1996; Binley et al. 1996; Poignard et al. 1996a; McDougal et al. 1996; Manca et al. 1995a; Sattentau et al. 1995; Chen et al. 1995; Laal et al. 1994; Spear et al. 1993; Eddleston et al. 1993; Sattentau & Moore 1991; Robinson et al. 1991; Xu et al. 1991; Gorny et al. 1989; Pinter et al. 1989;

**Keywords** antibody binding site definition and exposure, antibody interactions, complement, enhancing activity, immunotoxin, kinetics, mimotopes, review, subtype comparisons, vaccine antigen design, variant cross-recognition or cross-neutralization

 50-69: NIH AIDS Research and Reference Reagent Program: 531.

Till et al. 1989

- 50-69: This is one of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- 50-69: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of 17b and F105 was decreased by trypsin, but increased by thrombin. gp41 MAbs 246D, 98.6, 50-69, were decreased by trypsin, unaltered by thrombin, while NAb 2F5 binding was increased by thrombin. Thrombin may increase HIV-induced cell fusion in blood by causing a conformational activating shift in gp120. Ling et al. [2004] (antibody binding site definition and exposure)
- 50-69: Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. SF162 and each of the five glycosyation mutants studied were all neutralization resistant to 50-69. V3 glycans tended to shield V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans shielded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004] (antibody binding site definition and exposure, vaccine antigen design)
- 50-69: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I MAbs 50-69, F240, 240-D,3D6, and 246-D recognize a nonhelical hydrophobic region, positions 598-604, that forms a disulfide loop in the six-helix bundle. Cluster II MAbs 98-6 and 126-6 recognized residues 644-663 of gp41, a portion of the second heptad repeat. These MAbs were

found to behave similarly, so 50-69 and 98-6 were used as representatives. Exposure of cluster I and cluster II epitopes required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor. Binding to CD4 exposed hidden cluster I and II epitopes. The MAbs were found to bind to gp120/gp41 complexes, not to gp41 after shedding of gp120, and were localized to at fusing-cell interfaces. Kinetic and binding results indicate that these MAbs are exposed in transitional structures during the fusion process, possibly the prehairpin intermediate prior to co-receptor binding, although other intermediate structures may be involved. They do not bind once syncytia begin to show extensive cyotplasmic mixing. These MAbs failed to inhibit fusion. The NAb 2F5 has a very different behavior in this study. Finnegan *et al.* [2002] (antibody binding site definition and exposure, kinetics)

- 50-69: Called 50-69D. Alanine mutations were introduced into the N- and C-terminal alpha-helices of gp41 to destabilize interhelical packing interactions in order to study their inhibitory effect on viral infectivity. These mutations were shown to inhibit viral replication though affecting the conformational transition to the fusion-active form of gp41, and allow increased inhibition by gp41 peptides. 2F5 senstivity is increased in the mutated viruses, presumably because 2F5s neutralization activity is focused on the transition to the fusion active state. No other gp41 MAb against tested, including NC-1, 50-69D, 1281, 98-6D, 246-D and F240, neutralized the parental or the fusion-deficient mutated viruses. Follis *et al.* [2002] (antibody binding site definition and exposure)
- 50-69: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001] (antibody interactions)
- 50-69: This paper primarily concerns 4E10 and Z13, MAbs that both bind proximally to the 2F5 binding site to a conserved epitope, and that neutralize some primary isolates from clades B, C, and E MAb 50-69 binding to infected cells is enhanced by sCD4, while 4E10 and Z13 binding is essentially unaltered. Zwick *et al.* [2001b] (antibody binding site definition and exposure)
- 50-69: A cluster I epitope that binds to rgp41 567-647, recognizes a peptide N51-C43 complex trimer of heterodimers that approximates the core of the fusogenic form of gp41 this MAb doesn't react with either of the peptides N51 or C43 individually MAbs 50-69 and 1367 had similar properties MAb 50-69 bound the fusogenic form of the protein in liquid phase. Gorny & Zolla-Pazner [2000] (antibody binding site definition and exposure)
- 50-69: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, but gp41 MAb 50-69 bound with a 5 fold preference for the oligomer, while other gp41 MAbs (1367, 98-6, 167-D, 1281, 1342, and 1379) did not show a preference. Gorny *et al.* [2000] (antibody binding site definition and exposure)

- 50-69: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 cluster I anti-gp41 MAbs which showed good cross clade reactivity 50-69 bound the majority of isolates although binding was moderate to weak specifies discontinuous binding site range as aa 579-613. Nyambi et al. [2000] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 50-69: Mutations in BH10 gp160, W596Y and T605A, as well as deletions of 605-609 (TTAVP) and 597-609 (GCSGK-LICTTAVP), abrogate binding of enhancing MAbs 86, 240D, 50-69, and 246-D 5/6 enhancing MAbs identified to date bind to the immunodominant region 579-613 identifies noncontiguous W596-G597-C598 and C604-T605 as minimal epitope. Mitchell *et al.* [1998] (antibody binding site definition and exposure)
- 50-69: Abs that recognize discontinuous epitopes can identify mimotopes from a phage peptide display library 50-69 maps to an immunodominant domain in gp41 three groups of peptides were selected, one which seems most closely related to gp41 sequence peptide consensus is WGCxx(RK)(x n)LxC the analogous gp41 sequence WGCSGKLIC is present in most M group clades, except D with a common L to H substitution. Boots *et al.* [1997] (mimotopes)
- 50-69: Binding of anti-gp120 MAbs IgG1b12 or 654-30D does not mediate significant exposure of the gp41 epitopes for MAbs 2F5 and 50-69. Stamatatos *et al.* [1997] (antibody interactions)
- 50-69: Binds to a linear epitope located in the cluster I region binding of 50-69 and 240-D inhibited by Fabs A1, A4, M8B, M26B, M12B and T2. Binley *et al.* [1996] (antibody binding site definition and exposure)
- 50-69: Used to test exposure of gp41 upon sCD4 binding. Klasse & Sattentau [1996]
- 50-69: Does not neutralize HIV-1 LAI. McDougal *et al.* [1996] (variant cross-recognition or cross-neutralization)
- 50-69: Prebinding of anti-V3, and CD4i MAbs 48d and 17b, but not anti-V2 neutralizing MAbs, expose the 50-69 epitope. Poignard *et al.* [1996a] (antibody interactions)
- 50-69: One of several anti-gp41 MAbs that bind to a gp41-maltose binding fusion protein designed to study the leucine zipper domain of gp41, showing that the construct has retained aspects of normal gp41 conformation. Chen *et al.* [1995] (antibody binding site definition and exposure)
- 50-69: Virions complexed to gp41 Ab facilitate presentation of p66 RT epitopes to Th cells. Manca *et al.* [1995a]
- 50-69: Preferentially binds oligomer binding increased after pretreatment of infected cells with sCD4 – binding domain overlaps site that is critical for gp120-gp41 association. Sattentau et al. [1995] (antibody binding site definition and exposure)
- 50-69: Epitope described as cluster I, 601-604, conformational

   does not neutralize IIIB or synergize neutralization by anti-V3 MAb 447-52D or by CD4 BS MAbs. Laal *et al.* [1994] (antibody binding site definition and exposure, antibody interactions)
- 50-69: Called SZ-50.69 binds to an epitope within aa 579-613. Eddleston *et al.* [1993] (antibody binding site definition and exposure)
- 50-69: Did not mediate deposition of complement component C3 on HIV infected cells unless cells were pre-incubated with

- sCD4 complement mediated virolysis of MN and IIIB in the presence of sCD4. Spear *et al.* [1993] (**complement**)
- 50-69: Enhances HIV-1 infection *in vitro* synergizes with huMAb 120-16 *in vitro* to enhance HIV-1 infection to level approaching that found in polyclonal anti-HIV serum. Robinson *et al.* [1991] (antibody interactions, enhancing activity)
- 50-69: Two fold increase in binding to gp120 in the presence of bound sCD4. Sattentau & Moore [1991] (antibody binding site definition and exposure)
- 50-69: The epitope is affected by the conformation conferred by the two cysteines at amino acids 598 and 604. Xu *et al.* [1991] (antibody binding site definition and exposure)
- 50-69: Kills HIV-infected cells when coupled to deglycosylated ricin A chain. Gorny et al. [1989] (immunotoxin)
- 50-69: Reacts preferentially with gp160 oligomer, compared to gp41 monomer. Pinter *et al.* [1989] (antibody binding site definition and exposure)
- 50-69: Combined with deglycosylated A chain of ricin is toxic to lines of HIV-infected T cells (H9) and monocytes (U937). Till et al. [1989] (immunotoxin)

**No.** 663 **MAb ID** 9-11

HXB2 Location gp160 (579-604)

Author Location gp41 (584–609)

Epitope RILAVERYLKDQQLLGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: gp160

Species (Isotype) mouse (IgG1)

References Mani et al. 1994

• 9-11: required the C-C disulfide bridge and loop formation, can bind simultaneously with 41-1. Mani *et al.* [1994]

No. 664

**MAb ID** 98-43

HXB2 Location gp160 (579–604)

Author Location gp41 (579–604 HXB2)

Epitope RILAVERYLKDQQLLGIWGCSGKLIC

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG2\kappa$ )

**References** Xu *et al.* 1991; Tyler *et al.* 1990; Gorny *et al.* 1989; Pinter *et al.* 1989

- 98-43: NIH AIDS Research and Reference Reagent Program: 1241.
- 98-43: 579-604 binds in the immunodominant region. Xu et al. [1991]
- 98-43: Poor ADCC (in contrast to MAb 120-16, gp41(644-663)). Tyler et al. [1990]
- 98-43: Reacts equally well with oligomer and monomer. Pinter *et al.* [1989]

No. 665

**MAb ID** 41-1 (41.1)

HXB2 Location gp160 (579–608)

Author Location gp41 (584–609)

Epitope RILAVERYLKDQQLLGIWGCSGKLICTTAV

Neutralizing

Immunogen vaccine

*Vector/Type:* protein *HIV component:* gp160

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

**References** Pincus *et al.* 1998; Pincus *et al.* 1996; Mani *et al.* 1994; Pincus & McClure 1993; Pincus *et al.* 1991; Dalgleish *et al.* 1988; Gosting *et al.* 1987

- 41-1: Also called 41.1, although possibly not, the literature is confusing because two gp41 MAbs that bind to this region with similar names (dash versus period) are listed as murine and human.
- 41-1: Called 41.1, and described as a human MAb, binding 579-604 a panel of immunotoxins was generated by linking Env MAbs to ricin A immunotoxins mediated cell killing, but killing was not directly proportional to binding. Pincus *et al.* [1996]
- 41-1: This antibody to gp41(584-609) Mani *et al.* [1994] seems to have been named the same as a different MAb to gp41(735-752 IIIB) Dalgleish *et al.* [1988]. Dalgleish *et al.* [1988]; Mani *et al.* [1994]
- 41-1: Did not require the C-C disulfide bridge and loop formation, can bind simultaneously with 9-11. Mani *et al.* [1994]
- 41-1: Called 41.1, and described as a human MAb cross-competes with 41.4 sCD4 enhances the efficacy of immunotoxins *in vitro* 30-fold MAb was coupled to ricin A chain (RAC). Pincus & McClure [1993]
- 41-1: Efficacious as an immunotoxin when coupled to RAC gave linear epitope as gp160 579-603. Pincus *et al.* [1991]
- 41-1: This antibody seems to have been named the same as a different MAb to gp41(735-752). Dalgleish *et al.* [1988]
- 41-1: Broadly reactive. Gosting et al. [1987]

No. 666

**MAb ID** 41.4

**HXB2 Location** gp160 (579–608)

Author Location gp41 (584–609)

Epitope RILAVERYLKDQQLLGIWGCSGKLICTTAV

Neutralizing

Immunogen

Species (Isotype)

Research Contact Jan McClure, Bristol-Myers Squibb Pharmaceutical Res Inst, Seattle, WA

References Pincus & McClure 1993

• 41.4: Binds to peptide weakly, but to gp160 with higher affinity than 41.1, and cross-competes with 41.1 – probably conformational – MAb was coupled to ricin A chain (RAC) – sCD4 enhances the efficacy of immunotoxins *in vitro* 30-fold. Pincus & McClure [1993]

**No.** 667

MAb ID Fab A1 (A1)

HXB2 Location gp160 (579–608)

Author Location gp41 (584–609 LAI)

Epitope RILAVERYLKDQQLLGIWGCSGKLICTTAV

Subtype B

Neutralizing no

Immunogen HIV-1 infection **Species (Isotype)** human (IgG1 $\kappa$ )

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** anti-idiotype, antibody generation, antibody sequence, variable domain, review

- Fab A1: Called A1. One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- Fab A1: Binds to cluster I region competes with MAbs 240-D and 50-69 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (anti-idiotype, antibody generation, antibody sequence, variable domain)

No. 668

MAb ID Fab A4 (A4)

HXB2 Location gp160 (579-608)

Author Location gp41 (584-609 LAI)

Epitope RILAVERYLKDQQLLGIWGCSGKLICTTAV

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

- Fab A4: Called A4. One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- Fab A4: Binds to cluster I region competes with MAbs 240-D and 50-69 conformation sensitive variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

No. 669

**MAb ID** Fab M12B (M12B)

HXB2 Location gp160 (579–608)

**Author Location** gp41 (584–609 LAI)

Epitope RILAVERYLKDQQLLGIWGCSGKLICTTAV

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

- Fab M12B: Called M12B. One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- Fab M12B: Binds to cluster I region competes with MAbs 240-D and 50-69 conformation sensitive variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

**No.** 670

MAb ID Fab M26B (M26B)

HXB2 Location gp160 (579-608)

Author Location gp41 (584-609 LAI)

Epitope RILAVERYLKDQQLLGIWGCSGKLICTTAV

Subtype B Neutralizing no

Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ )

> References Gorny & Zolla-Pazner 2004; Binley et al. 1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

- Fab M26B: Called M26B. One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- Fab M26B: Binds to cluster I region competes with MAbs 240-D and 50-69 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

No. 671

MAb ID Fab M8B (M8B)

HXB2 Location gp160 (579-608)

Author Location gp41 (584–609 LAI)

Epitope RILAVERYLKDQQLLGIWGCSGKLICTTAV

Subtype B Neutralizing no

Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ )

> References Gorny & Zolla-Pazner 2004; Binley et al. 1996

Keywords antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

- Fab M8B: Called M8B. One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- Fab M8B: Binds to cluster I region competes with MAbs 240-D and 50-69 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

No. 672

MAb ID Fab T2 (T2)

HXB2 Location gp160 (579-608)

Author Location gp41 (584–609 LAI)

Epitope RILAVERYLKDQQLLGIWGCSGKLICTTAV

Subtype B Neutralizing no

Immunogen HIV-1 infection **Species (Isotype)** human (IgG1 $\kappa$ )

References Gorny & Zolla-Pazner 2004; Binley et al.

1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

- Fab T2: Called T2. One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- Fab T2: Binds to cluster I region competes with MAbs 240-D and 50-69 - conformation sensitive - variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

No. 673

MAb ID polyclonal

HXB2 Location gp160 (579–608)

Author Location gp41

Epitope RVAVERYLKDQQLLGIWGCSGKICTTAV

Subtype D, multiple

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

References Barin et al. 2005

Country France

Keywords acute/early infection, assay development

· A combination of 4 antigenic regions was used to differentiate between early (<180 days) and chronic infection. These regions were: p24; the gp41 peptide spanning the immunodominant epitope (IDE) of gp41, RVAVERYLKDQQLLGIWGC-SGKICTTAV, and a subtype D version of this peptide; 5 V3 consensus peptides including A, B, C, D, and CRF01-AE; and Integrase. V3 and the IDE provide the best discrimination, with >20 fold higher levels in chronic infection when assayed by EIA using dried serum spots. Antibodies to Integrase and p24 were not as distinctive, and people tend to lose, not increase, responses to p24 over time. This assay can be used to identify samples from early infection with high sensitivity and specificity. Barin et al. [2005] (assay development, acute/early infection)

No. 674

**MAb ID** 86 (No. 86)

HXB2 Location gp160 (579–613)

Author Location gp41 (586–620 IIIB)

Epitope RILAVERYLKDQQLLGIWGCSGKLICTTAVPW-

NAS

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Research Contact Evan Hersh and Yoh-Ichi Matsumoto

References Gorny & Zolla-Pazner 2004; Mitchell et al. 1998; Wisnewski et al. 1996; Moran et al. 1993; Pincus et al. 1991; Robinson et al. 1990c; Robinson et al. 1990b; Sugano et al.

Keywords antibody binding site definition and exposure, antibody sequence, variable domain, complement, enhancing activity, immunotoxin, review, variant cross-recognition or cross-

neutralization

- 86: NIH AIDS Research and Reference Reagent Program: 380. V10-9: One of 24 MAbs and Fabs in this database that bind to
- 86: This is one of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- 86: Mutations in BH10 gp160, W596Y and T605A, as well as deletions of 605-609 (TTAVP) and 597-609 (GCSGK-LICTTAVP), abrogate binding of enhancing MAbs 86, 240D, 50-69, and 246-D 5/6 enhancing MAbs identified to date bind to the immunodominant region 579-613. Mitchell *et al.* [1998] (enhancing activity, variant cross-recognition or cross-neutralization)
- 86: 86 is V H1 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski *et al.* [1996] (antibody sequence, variable domain)
- 86: Heavy (V HI) and light (V kappaI) chain sequenced enhancing activity similar germline sequence to MAb S1-1, but very different activity. Moran *et al.* [1993] (enhancing activity, antibody sequence, variable domain)
- 86: Poor immunotoxin activity when coupled to RAC peptide binding stated to be aa 579-603. Pincus *et al.* [1991] (**antibody binding site definition and exposure**, **immunotoxin**)
- 86: Antibody dependent enhancement (ADE) of HIV-1 IIIB infectivity in the presence of complement. Robinson *et al.* [1990b] (**complement, enhancing activity**)
- 86: Peptide 586-620 blocks complement mediated ADE. Robinson et al. [1990c] (enhancing activity)
- 86: Reacts with gp41 and also reacted weakly with gp120.
   Sugano et al. [1988] (antibody binding site definition and exposure)

No. 675

MAb ID polyclonal

HXB2 Location gp160 (580-597)

Author Location gp41 (584–602)

Epitope ILAVERYLKDQQLLGIWG

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human

References Petrov et al. 1990

 Immunodominant and broadly reactive peptide. Petrov et al. [1990]

**No.** 676

**MAb ID** V10-9

**HXB2 Location** gp160 (580–613)

Author Location gp41 (586-620 IIIB)

Epitope ILAVERYLKDQQLLGIWGCSGKLICTTAVPWN-

AS

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG1 $\kappa$ )

References Gorny & Zolla-Pazner 2004; Robinson et al.

1990c; Robinson et al. 1990b

**Keywords** antibody interactions, enhancing activity, review

 V10-9: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604).
 Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)

- V10-9: Antibody dependent enhancement (ADE) of HIV-1 IIIB infectivity, synergistically enhanced by MAb 120-16. Robinson *et al.* [1990b] (antibody interactions, enhancing activity)
- V10-9: Peptide 586-620 blocks complement mediated ADE. Robinson *et al.* [1990c] (**enhancing activity**)

No. 677

MAb ID polyclonal

HXB2 Location gp160 (582-589)

Author Location gp41 (589-596)

Epitope AVERYLKD

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

References Klasse et al. 1991

Substitutions and deletions in peptide 583-599 were systematically studied – alterations in AVERYLKD abrogated the antigenicity of peptides with most of 14 human sera. Klasse et al. [1991]

No. 678

MAb ID anti-P1

HXB2 Location gp160 (582–592)

Author Location gp41 (579-613)

Epitope AVERYLKDQQL

Subtype B

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype)

References Ferraz et al. 2004

Keywords assay development

The B-cell epitope of P1 was incorporated into the solvent-exposed loop of the E. coli betagalactosidase enzyme for use as an analytical biosensor to permit enzyme substrate analysis to better understand the conversion of conformational stimulous into enzymatic signal. Ferraz et al. [2004] (assay development)

**No.** 679

MAb ID polyclonal

HXB2 Location gp160 (584–604)

Author Location gp41 (74–94)

Epitope ERYLKDQLLGIWGCSGKLIC

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Shafferman et al. 1989

• Immunogenic domain useful for diagnostics. Shafferman *et al.* [1989]

**No.** 680

MAb ID polyclonal

**HXB2 Location** gp160 (584–612)

Author Location gp41 (587–617 BRU)

Epitope ERYLKDQQLLGIWGCSGKLICTTAVPWNA

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human

References Hernandez et al. 2000

 Chimeric peptide combining two peptides gp160(495-516 and 584-612) served as a specific and broadly reactive antigen for diagnostic detection of HIV-1. Hernandez *et al.* [2000]

**No.** 681

MAb ID 2F11

**HXB2 Location** gp160 (589–600)

Author Location gp41 (589–600 HXB2)

Epitope DQQLLGIWGCSG

Subtype B

Neutralizing no

Immunogen HIV-1 infection Species (Isotype) human (IgG1)

**References** Gorny & Zolla-Pazner 2004; Enshell-Seijffers

et al. 2001; Eaton et al. 1994

Keywords ADCC, enhancing activity, review

- 2F11: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- 2F11: Monoclonal antibodies to this epitope have distinct phenotypes—41-7 and 1B8.env were found to be not neutralizing, 2F11 possibly enhancing, and clone 3 beneficial. Enshell-Seijffers *et al.* [2001] (**enhancing activity**)
- 2F11: Enhances infectivity even in the absence of complement does not mediate ADCC or neutralize virus. Eaton *et al.* [1994]
   (ADCC, enhancing activity)

No. 682

MAb ID 246-D (SZ-246.D, 246, 246D)

**HXB2 Location** gp160 (590–597)

Author Location gp41 (579–604 HXB2)

Epitope qqLLGIWg

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**Ab Type** gp41 cluster I

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu), NYU Med

iasul@incicio.ined.nyu), N10 Med

Center, NY, NY

References Ling et al. 2004; Gorny & Zolla-Pazner 2004;

Finnegan et al. 2002; Follis et al. 2002; Edwards et al. 2002; Gorny et al. 2002; Verrier et al. 2001; Nyambi et al. 2000; Gorny & Zolla-Pazner 2000; Mitchell et al. 1998; Hioe et al. 1997b; Earl et al. 1997; Saarloos et al. 1995; Manca et al. 1995; Forthal et al. 1995; Eddleston et al. 1993; Spear et al. 1993;

Robinson et al. 1991; Xu et al. 1991

**Keywords** antibody binding site definition and exposure, antibody interactions, complement, enhancing activity, kinetics, review, subtype comparisons, variant cross-recognition or cross-

neutralization

 246-D: NIH AIDS Research and Reference Reagent Program: 1245.

- 246-D: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- 246-D: Called 246D. The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of 17b and F105 was decreased by trypsin, but increased by thrombin. gp41 MAbs 246D, 98.6, 50-69, were decreased by trypsin, unaltered by thrombin, while NAb 2F5 binding was increased by thrombin. Thrombin may increase HIV-induced cell fusion in blood by causing a conformational activating shift in gp120. Ling et al. [2004] (antibody binding site definition and exposure)
- 246-D: Called 246D Truncation of the gp41 cytoplasmic domain of X4, R5, and X4R5 viruses forces a conformation that more closely resembles the CD4 bound state of the external Envelope, enhancing binding of CD4i MAbs 17b and 48d and of CD4BS MAbs F105, b12, and in most cases of glycosylation site dependent MAb 2G12 and the anti-gp41 MAb 246D in contrast, binding of the anti-V2 MAb 697D and the anti-V3 MAb 694/98D were not affected viruses bearing the truncation were more sensitive to neutralization by MAbs 48d, b12, and 2G12 the anti-C5 MAb 1331A was used to track levels of cell surface expression of the mutated proteins. Edwards *et al.* [2002] (antibody binding site definition and exposure)
- 246-D: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I MAbs 50-69, F240, 240-D,3D6, and 246-D recognize a nonhelical hydrophobic region, positions 598-604, that forms a disulfide loop in the six-helix bundle. Cluster II MAbs 98-6 and 126-6 recognized residues 644-663 of gp41, a portion of the second heptad repeat. These MAbs were found to behave similarly, so 50-69 and 98-6 were used as representatives. Exposure of cluster I and cluster II epitopes required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor. Binding to CD4 exposed hidden cluster I and II epitopes. The MAbs were found to bind to gp120/gp41 complexes, not to gp41 after shedding of gp120, and were localized to at fusing-cell interfaces. Kinetic and binding results indicate that these MAbs are exposed in transitional structures during the fusion process, possibly the prehairpin intermediate prior to co-receptor binding, although other intermediate structures may be involved. They do not bind once syncytia begin to show extensive cyotplasmic mixing. These MAbs failed to inhibit fusion. The NAb 2F5 has a very different behavior in this study. Finnegan et al. [2002] (antibody binding site definition and exposure, kinetics)
- 246-D: Alanine mutations were introduced into the N- and C-terminal alpha-helices of gp41 to destabilize interhelical packing interactions in order to study their inhibitory effect on viral infectivity. These mutations were shown to inhibit viral replication though affecting the conformational transition to the fusion-active form of gp41, and allow increased inhibition

by gp41 peptides. 2F5 senstivity is increased in the mutated viruses, presumably because 2F5s neutralization activity is focused on the transition to the fusion active state. No other gp41 MAb against tested, including NC-1, 50-69D, 1281, 98-6D, 246-D and F240, neutralized the parental or the fusion-deficient mutated viruses. Follis *et al.* [2002] (antibody binding site definition and exposure)

- 246-D: Called 246 Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated – the six new MAbs all bind to the tip of the V3 loop and cross-compete with the MAb 447-52D and are conformationally sensitive – MAbs showed cross-clade binding to native, intact virions and the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay – five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization studies), 654 (anti-CD4BS used as a conformation-sensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control), and MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades tested, A, B, C, D, F and CRF01 (clade E). Gorny et al. [2002] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 246-D: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001] (antibody interactions)
- 246-D: Core epitope aa 591 to 597, a cluster I epitope that does
  not bind to either a peptide complex that approximates the core
  of the fusogenic form of gp41 or the individual peptides N51
  and C43 that form this structure MAbs 181-D and 246-D had
  similar properties. Gorny & Zolla-Pazner [2000] (antibody
  binding site definition and exposure)
- 246-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 cluster I anti-gp41 MAbs which showed good cross clade reactivity 246-D bound strongly or moderately to all 26 HIV-1 group M clades viruses tested and showed the strongest binding of all anti-Env MAbs tested, including the V3 and C5 region MAbs notes core epitope as LLGI no neutralizing activity was observed when 246-D was tested with five isolates. Nyambi *et al.* [2000] (subtype comparisons)
- 246-D: Mutations in BH10 gp160, W596Y and T605A, as well as deletions of 605-609 (TTAVP) and 597-609 (GCSGKLICT-TAVP), abrogate binding of enhancing MAbs 86, 240D, 50-69, and 246-D 5/6 enhancing MAbs identified to date bind to the immunodominant region 579-613. Mitchell *et al.* [1998] (antibody binding site definition and exposure)
- 246-D: This antibody, along with murine MAb D61, can be blocked by any of a group of 8 conformational MAbs (M10, D41, D54, T4, T6, T9, T10 and T35). Earl *et al.* [1997] (antibody interactions)
- 246-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs
   BZ167 was the only isolate inhibited by all polyclonal sera

and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) – isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations and 246-D neutralized 91US056 – US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe *et al.* [1997b] (variant cross-recognition or cross-neutralization)

- 246-D: No neutralizing activity, both ADCC and viral enhancing activity. Forthal *et al.* [1995] (complement, enhancing activity)
- 246-D: Virions complexed to gp41 Ab facilitate presentation of p66 RT epitopes to Th cells. Manca *et al.* [1995a]
- 246-D: Ab-mediated activation of complement on HIV+ cells is higher than Ab independent activation—what has been termed "Ab independent" in fact results in part from IgM in normal human serum that is HIV-cross-reactive. Saarloos *et al.* [1995] (**complement**)
- 246-D: Called SZ-246.D. Eddleston et al. [1993]
- 246-D: Did not mediate deposition of complement component C3 on HIV infected cells unless cells were pre-incubated with sCD4. Spear *et al.* [1993] (complement)
- 246-D: No neutralizing activity, some enhancing activity.
   Robinson et al. [1991] (enhancing activity)
- 246-D: Fine mapping indicates core is LLGI. Xu *et al.* [1991] (antibody binding site definition and exposure)

**No.** 683

MAb ID polyclonal

HXB2 Location gp160 (590–607)

**Author Location** gp41

Epitope QQLLGIWGCSGKLICTTA

Subtype B, CRF01\_AE

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Parekh et al. 2002

A simple enzyme immunoassay (EIA) that detects increasing levels of anti-HIV IgG after seroconversion can be used for detecting recent HIV-1 infection – longitudinal specimens from 139 incident infections in the US and Thailand were used in the study – the method was generally applicable for HIV-1 subtypes A, B, C, D and E(CRF01). Parekh *et al.* [2002]

No. 684

MAb ID 9G5A

**HXB2 Location** gp160 (591–594)

Author Location gp41 (596–599 IIIB)

**Epitope** QLLG

**Neutralizing** 

Immunogen anti-idiotype

Species (Isotype) mouse (IgM)

**References** Beretta & Dalgleish 1994; Lopalco *et al.* 1993 9G5A: Anti-idiotype to gp120 C terminus (C5 region) MAb

M38. Lopalco *et al.* [1993]

**No.** 685

**MAb ID** 181-D (SZ-181.D)

HXB2 Location gp160 (591-597)

Author Location gp41 (591–597 HXB2)

Epitope qLLGIWg

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG2\kappa$ )

Ab Type gp41 cluster I

Research Contact Susan Zolla-Pazner (Zolla-Pazner NVII NV

las01@mcrcr6.med.nyu), NYU, NY

References Gorny & Zolla-Pazner 2004; Nyambi et al. 2000; Gorny & Zolla-Pazner 2000; Fontenot et al. 1995; Forthal et al. 1995; Eddleston et al. 1993; Robinson et al. 1991; Xu et al.

**Keywords** ADCC, antibody binding site definition and exposure, enhancing activity, review, subtype comparisons, variant cross-recognition or cross-neutralization

- 181-D: This is one of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- 181-D: Core epitope aa 591 to 597, a cluster I epitope that does
  not bind to either a peptide complex that approximates the core
  of the fusogenic form of gp41 or the individual peptides N51
  and C43 that form this structure MAbs 181-D and 246-D had
  similar properties. Gorny & Zolla-Pazner [2000] (antibody
  binding site definition and exposure)
- 181-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 cluster I anti-gp41 MAbs which showed good cross clade reactivity 181-D bound the majority of isolates although binding was moderate to weak. Nyambi et al. [2000] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 181-D: No neutralizing, no ADCC, and no viral enhancing activity. Forthal *et al.* [1995] (ADCC, enhancing activity)
- 181-D: Called SZ-181.D. Eddleston et al. [1993]
- 181-D: No enhancing or neutralization activity. Robinson *et al.* [1991] (**enhancing activity**)
- 181-D: Fine mapping indicates core is LLGIW. Xu *et al.* [1991] (antibody binding site definition and exposure)

**No.** 686

**MAb ID** 240-D (F240)

HXB2 Location gp160 (592–600)

Author Location gp41 (592–600 HXB2)

Epitope LLGIWGCSG

Subtype B

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp41 cluster I

Research Contact Susan Zolla-Pazner

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(Zol-

**References** Gorny & Zolla-Pazner 2004; Finnegan *et al.* 2002; Nyambi *et al.* 2000; Mitchell *et al.* 1998; Wisnewski *et al.* 1996; Wisnewski *et al.* 1995; Binley *et al.* 1996; Spear *et al.* 1993; Robinson *et al.* 1991; Xu *et al.* 1991

**Keywords** antibody binding site definition and exposure, antibody sequence, variable domain, complement, enhancing activity, kinetics, review, subtype comparisons, variant cross-recognition or cross-neutralization

- (Zol- 240-D: NIH AIDS Research and Reference Reagent Program: 1242.
  - 240-D: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604).
     Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
  - 240-D: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I MAbs 50-69, F240, 240-D,3D6, and 246-D recognize a nonhelical hydrophobic region, positions 598-604, that forms a disulfide loop in the six-helix bundle. Cluster II MAbs 98-6 and 126-6 recognized residues 644-663 of gp41, a portion of the second heptad repeat. These MAbs were found to behave similarly, so 50-69 and 98-6 were used as representatives. Exposure of cluster I and cluster II epitopes required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor. Binding to CD4 exposed hidden cluster I and II epitopes. The MAbs were found to bind to gp120/gp41 complexes, not to gp41 after shedding of gp120, and were localized to at fusing-cell interfaces. Kinetic and binding results indicate that these MAbs are exposed in transitional structures during the fusion process, possibly the prehairpin intermediate prior to co-receptor binding, although other intermediate structures may be involved. They do not bind once syncytia begin to show extensive cyotplasmic mixing. These MAbs failed to inhibit fusion. The NAb 2F5 has a very different behavior in this study. Finnegan et al. [2002] (antibody binding site definition and exposure, kinetics)
  - 240-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 cluster I anti-gp41 MAbs which showed good cross clade reactivity 246-D bound strongly or moderately to 24/26 HIV-1 group M clades viruses tested. Nyambi *et al.* [2000] (variant cross-recognition or cross-neutralization, subtype comparisons)
  - 240-D: Mutations in BH10 gp160, W596Y and T605A, as well as deletions of 605-609 (TTAVP) and 597-609 (GCSGKLICT-TAVP), abrogate binding of enhancing MAbs 86, 240D, 50-69, and 246-D 5/6 enhancing MAbs identified to date bind to the immunodominant region 579-613. Mitchell *et al.* [1998] (**enhancing activity**)
  - 240-D: Binds to a linear epitope located in the cluster I region binding of 50-69 and 240-D inhibited by Fabs A1, A4, M8B, M26B, M12B and T2. Binley *et al.* [1996] (antibody binding site definition and exposure)
  - 240-D: V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski *et al.* [1996] (antibody sequence, variable domain)

C3 on HIV infected cells. Spear *et al.* [1993] (**complement**)

- 240-D: No neutralizing activity, some enhancing activity. Robinson et al. [1991] (enhancing activity)
- 240-D: Fine mapping indicates core is IWG. Xu et al. [1991] (antibody binding site definition and exposure)

No. 687 MAb ID F240

HXB2 Location gp160 (592-606)

Author Location gp41 (592–606 BH10)

Epitope LLGIWGCSGKLICTT

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type gp41 cluster I

Research Contact L. Cavacina or M. Posner, Dept. of Med. Har-

vard Med. School, Boston MA, USA

Cavacini et al. 2003; Cavacini et al. 2002;

References Liu et al. 2005a; Gorny & Zolla-Pazner 2004; Finnegan et al. 2002; Follis et al. 2002;

York et al. 2001; Cavacini et al. 1998a

**Keywords** antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain, co-receptor, enhancing activity, review, variant cross-recognition or crossneutralization

- F240: Transduction of human CD4+ H9 T cells with both the intracellularly expressed and secreted forms of the single-chain F240 Ab inhibited MN virus production. The secreted form was more potent. Viral replication of HIV-1 primary isolates was not reduced. Liu et al. [2005a]
- F240: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- F240: The MAb B4e8 binds to the base of the V3 loop, neutralizes multiple primary isolates and was studied for interaction with other MAbs. Anti-gp41 MAb F240 could inhibit B4e8 neutralization. Cavacini et al. [2003] (antibody interactions)
- F240: This study examined antibody interactions, binding and neutralization with a B clade R5 isolate (92US660) and R5X4 isolate (92HT593). Abs generally bound and neutralized the R5X4 isolate better than the R5 isolate, with the exception of F240 which bound both equally well, which captured more virus than any other human MAb tested, and didn't neutralize either isolate.F240 enhanced the binding of CD4BS MAbs IgG1b12 and F105 and the gp41 MAb 2F5 for both R5X4 and R5 isolates. F240 binding to gp41 was not affected by the binding of the V3 loop MAb B4a1, but preincubation with F240 could enhance B4a1 binding of the R5 isolate. Synergistic neutralization between F240 and CD4i MAbs 17b and 48d was noted for the R5X4 but not the R5 isolate, and F240 also enhanced neutralization of the R5X4 isolate by 2F5, but had no effect on R5 virus. In contrast, F240 combined with 2G12 demonstrated enhanced neutralization of R5 virus at low Ab concentrations. Cavacini et al. [2002] (antibody interactions, co-receptor)

- 240-D: Did not mediate deposition of complement component F240: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I MAbs 50-69, F240, 240-D,3D6, and 246-D recognize a nonhelical hydrophobic region, positions 598-604, that forms a disulfide loop in the six-helix bundle. Cluster II MAbs 98-6 and 126-6 recognized residues 644-663 of gp41, a portion of the second heptad repeat. These MAbs were found to behave similarly, so 50-69 and 98-6 were used as representatives. Exposure of cluster I and cluster II epitopes required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor. Binding to CD4 exposed hidden cluster I and II epitopes. The MAbs were found to bind to gp120/gp41 complexes, not to gp41 after shedding of gp120, and were localized to at fusing-cell interfaces. Kinetic and binding results indicate that these MAbs are exposed in transitional structures during the fusion process, possibly the prehairpin intermediate prior to co-receptor binding, although other intermediate structures may be involved. They do not bind once syncytia begin to show extensive cytoplasmic mixing. These MAbs failed to inhibit fusion. The NAb 2F5 has a very different behavior in this study. Finnegan et al. [2002] (antibody binding site definition and exposure)
  - F240: Alanine mutations were introduced into the N- and C-terminal alpha-helices of gp41 to destabilize interhelical packing interactions in order to study their inhibitory effect on viral infectivity. These mutations were shown to inhibit viral replication though affecting the conformational transition to the fusion-active form of gp41, and allow increased inhibition by gp41 peptides. 2F5 sensitivity is increased in the mutated viruses, presumably because 2F5s neutralization activity is focused on the transition to the fusion active state. No other gp41 MAb against tested, including NC-1, 50-69D, 1281, 98-6D, 246-D and F240, neutralized the parental or the fusion-deficient mutated viruses. Follis et al. [2002] (antibody binding site definition and exposure)
  - F240: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding. York et al. [2001] (variant cross-recognition or crossneutralization)
  - F240: Distinct from MAb 240-D, an antibody with a similar epitope in the immunodominant region of gp41 - dosedependent reactivity with HIV isolates RF, SF2, IIIB, and MN was observed – F240 had no neutralizing activity and enhances infection in the presence of complement - reactivity of F240 is enhanced by preincubation of cells with sCD4 or anti-CD4BS MAb F105 – heavy and light chain variable domains were sequenced, and a strong homology to hu MAb 3D6 was observed, as 3D6 binds to the same epitope, these MAbs may define a human Ab clonotype. Cavacini et al. [1998a] (enhancing activity, variant cross-recognition or cross-neutralization, antibody sequence, variable domain)

No. 688 MAb ID D49

**HXB2 Location** gp160 (592–608) **Author Location** gp41 (597–613)

Epitope LLGIWGCSGKLICTTAV

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: dimeric Env

Species (Isotype) mouse

Ab Type gp41 cluster I

References Earl et al. 1997; Earl et al. 1994

• D49: Binding maps to region 597-613: WGCSGKLICTTAVP-WNA – immunodominant region containing two Cys residues. Earl *et al.* [1997]

D49: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

No. 689

MAb ID D61

HXB2 Location gp160 (592–608)

Author Location gp41 (592-608 HXB2)

Epitope LLGIWGCSGKLICTTAV

Subtype B Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component:

dimeric Env

Species (Isotype) mouse

Ab Type gp41 cluster I

Research Contact Patricia Earl and Christopher Broder, NIH

**References** Golding *et al.* 2002b; Earl *et al.* 1997; Weissenhorn *et al.* 1996; Richardson *et al.* 1996;

Earl et al. 1994

Keywords antibody binding site definition and exposure,

antibody generation

- D61: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion the preincubation 31.5 C step did not alter the inhibitory activity of neutralizing Abs anti-gp41 2F5, or anti-gp120 2G12, IG1b12, 48d, and 17b nor did it alter two gp41 MAbs, T9 and D61, inability to inhibit fusion. Golding *et al.* [2002b] (antibody binding site definition and exposure)
- D61: Binding maps to region 597-613: WGCSGKLICTTAVP-WNA immunodominant region containing two Cys residues this antibody, along with human MAb 246-D, can be blocked by any of a group of 8 conformational MAbs (M10, D41, D54, T4, T6, T9, T10 and T35) members of this competition group are blocked by sera from HIV-1 + individuals. Earl *et al.* [1997] (antibody binding site definition and exposure)
- D61: Linear gp41 epitope in the cluster I region human sera blocked binding in oligomeric ELISA assay to a similar extent for gp41 MAbs D20, D43, D61, and T4. Richardson *et al.* [1996] (antibody binding site definition and exposure)

• D61: Does not precipitate gp41(21-166), but due to a structural difference in the disulfide bonding region near the two cysteines – the authors propose that this region may change conformation during the activation of the membrane fusion state of the HIV-1 glycoprotein. Weissenhorn *et al.* [1996] (antibody binding site definition and exposure)

D61: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994] (antibody generation)

**No.** 690

MAb ID T32

**HXB2 Location** gp160 (592–608)

Author Location gp41 (597-613)

Epitope LLGIWGCSGKLICTTAV

**Neutralizing** 

Immunogen vaccine

Vector/Type: tetrameric Env HIV compo-

nent: Env

Species (Isotype) mouse

Ab Type gp41 cluster I

Research Contact Patricia Earl and Christopher Broder, NIH

References Earl et al. 1997; Earl et al. 1994

- T32: Binding maps to region 597-613: WGCSGKLICTTAVP-WNA immunodominant region containing two Cys residues. Earl *et al.* [1997]
- T32: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

No. 691

MAb ID T34

**HXB2 Location** gp160 (592–608)

Author Location gp41 (597–613)

Epitope LLGIWGCSGKLICTTAV

Neutralizing

Immunogen vaccine

Vector/Type: tetrameric Env HIV compo-

nent: Env

Species (Isotype) mouse

Ab Type gp41 cluster I

Research Contact Patricia Earl and Christopher Broder, NIH

References Earl et al. 1997; Earl et al. 1994

- T34: Binding maps to region 597-613: WGCSGKLICTTAVPWNA immunodominant region containing two Cys residues. Earl *et al.* [1997]
- T34: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response an oligomer with no gp120/gp41 cleavage site was used as the immunogen. Earl *et al.* [1994]

No. 692

**MAb ID** 115.8

**HXB2 Location** gp160 (593–604)

**Author Location** gp41 (598–609)

Epitope LGLIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgM)

References Oldstone et al. 1991

• 115.8: Stimulated by immunization with the peptide: LGLI-WGCSGKLIC (aa 598-609) – poor reactivity with CSGKLIC – reacts well with longer HIV-2 peptide NSWGCAFRQVC as well as CAFRQVC – disulfide bond between cysteines required. Oldstone *et al.* [1991]

No. 693

MAb ID M-1

HXB2 Location gp160 (593-604)

Author Location gp41 (598-609)

Epitope LGIWGCSGKLIC

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

 $\textbf{Species (Isotype)} \ \ mouse \ (IgG1, IgG2b)$ 

References Yamada et al. 1991

• M-1: Unlike M-22, did not react to 43-kd protein found in rat and human astrocytes. Yamada *et al.* [1991]

**No.** 694

MAb ID M-11

HXB2 Location gp160 (593-604)

Author Location gp41 (598–609)

Epitope LGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgG1)

References Yamada et al. 1991

• M-11: Strongly reacted with a cellular 43-kd protein found in rat and human astrocytes as well as with gp41. Yamada *et al.* [1991]

No. 695

**MAb ID** M-13

HXB2 Location gp160 (593–604)

Author Location gp41 (598-609)

Epitope LGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgG2b)

References Yamada et al. 1991

• M-13: Reacted with a cellular 43-kd protein found in rat and human astrocytes as well as with gp41. Yamada *et al.* [1991]

No. 696

MAb ID M-2

**HXB2 Location** gp160 (593–604)

Author Location gp41 (598–609)

Epitope LGIWGCSGKLIC

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgG2b)

References Yamada et al. 1991

• M-2: Strongly reacted with a cellular 43-kd protein found in rat and human astrocytes as well as with gp41. Yamada *et al.* [1991]

**No.** 697

MAb ID M-22

HXB2 Location gp160 (593–604)

**Author Location** gp41 (598–609)

Epitope LGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgG2b)

References Yamada et al. 1991

• M-22: Strongest reaction of 12 anti-HIV-1 gp41 MAbs to a cellular 43-kd protein found in rat and human astrocytes. Yamada *et al.* [1991]

No. 698

MAb ID M-24

HXB2 Location gp160 (593-604)

Author Location gp41 (598–609)

Epitope LGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

**Species** (**Isotype**) mouse (IgG1)

References Yamada et al. 1991

• M-24: Strongly reacted with a cellular 43-kd protein found in rat and human astrocytes as well as with gp41. Yamada *et al.* [1991]

No. 699

MAb ID M-25

HXB2 Location gp160 (593-604)

Author Location gp41 (598–609)

Epitope LGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

**Species (Isotype)** mouse (IgG1)

References Yamada et al. 1991

• M-25: Reacted with a cellular 43-kd protein found in rat and human astrocytes as well as with gp41. Yamada *et al.* [1991]

**No.** 700

MAb ID M-28

HXB2 Location gp160 (593-604)

Author Location gp41 (598–609)

Epitope LGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgG1)

References Yamada et al. 1991

• M-28: Strongly reacted with a cellular 43-kd protein found in rat and human astrocytes as well as with gp41. Yamada *et al.* [1991]

**No.** 701

MAb ID M-29

HXB2 Location gp160 (593–604)

Author Location gp41 (598-609)

**Epitope** LGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgG1)

References Yamada et al. 1991

• M-29: Unlike M-22, did not react to 43-kd protein found in rat and human astrocytes. Yamada *et al.* [1991]

**No.** 702

MAb ID M-36

HXB2 Location gp160 (593-604)

Author Location gp41 (598–609)

Epitope LGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgG1)

References Yamada et al. 1991

• M-36: Unlike M-22, did not react to 43-kd protein found in rat and human astrocytes. Yamada *et al.* [1991]

**No.** 703

MAb ID M-4

HXB2 Location gp160 (593–604)

Author Location gp41 (598-609)

Epitope LGIWGCSGKLIC

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

**Species (Isotype)** mouse (IgG2b)

References Yamada et al. 1991

• M-4: Unlike M-22, did not react to 43-kd protein found in rat and human astrocytes. Yamada *et al.* [1991]

No. 704

MAb ID M-6

HXB2 Location gp160 (593-604)

Author Location gp41 (598-609)

Epitope LGIWGCSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgG2b)

References Yamada et al. 1991

• M-6: Unlike M-22, did not react to 43-kd protein found in rat and human astrocytes. Yamada *et al.* [1991]

No. 705

MAb ID polyclonal α598-609

HXB2 Location gp160 (594-601)

Author Location gp41 (598–609)

Epitope GIWGCSGK

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Poumbourios et al. 1992

alpha(598-609): Affinity purified from HIV-1 + plasma – immunodominant region, binds oligomer and monomer. Poumbourios et al. [1992]

**No.** 706

**MAb ID** 1B8.env (1B8)

**HXB2 Location** gp160 (594–604)

Author Location gp41 (594–605 HXB2)

Epitope GIWGCSGKLIC

Subtype B

Neutralizing no

**Immunogen** HIV-1 infection **Species** (**Isotype**) human ( $IgG2\lambda$ )

References Gorny & Zolla-Pazner 2004; Enshell-Seijffers

et al. 2001; Banapour et al. 1987

**Keywords** antibody binding site definition and exposure,

review, variant cross-recognition or cross-

neutralization

• 1B8B.env: Called 1B8. There are 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)

- 1B8.env: Monoclonal antibodies to this epitope have distinct phenotypes—41-7 and 1B8.env were found to be not neutralizing, 2F11 possibly enhancing, and clone 3 beneficial. Enshell-Seijffers *et al.* [2001]
- 1B8.env: Highly conserved epitope recognized by the majority of HIV-1 infected people MAb does not neutralize. Banapour *et al.* [1987] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)

No. 707

MAb ID polyclonal

**HXB2 Location** gp160 (594–609)

Author Location gp41 (601–616)

Epitope GIWGCSGKLICTTAVP

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human

References Petrov et al. 1990

 Immunodominant and broadly reactive peptide. Petrov et al. [1990]

No. 708

MAb ID polyclonal

HXB2 Location gp160 (595–607)

Author Location gp41 (600–612)

Epitope IWGCSGKLICTTA

Subtype B

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Belliard et al. 2003

**Country** France

Keywords rate of progression

• Sera from 101 slow progressors and 42 fast progressors were tested for responses to Tat peptides, and compared to responses to gp41 peptide 600-612, as anti-Tat antibodies had been shown by others to be elevated in slow progressors. Most patient sera react with this peptide, it is used in diagnostics. In this study, overall levels of Tat antibodies were not different in the two groups, however relative levels of antibodies to different Tat peptides and to this gp41 peptide were observed. Belliard *et al.* [2003] (**rate of progression**)

**No.** 709

MAb ID clone 3 (CL3)

HXB2 Location gp160 (597-606)

Author Location gp41 (597–606)

**Epitope** GCSGKLICTT

Subtype B

**Neutralizing** LP

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Research Contact BioClonetics (Philadelphia)

2004; Enshell-Seijffers et al. 2001; Cotropia et al. 1996; Cotropia et al. 1992; Broliden

References Ferrantelli et al. 2004a; Gorny & Zolla-Pazner

et al. 1989

**Keywords** antibody binding site definition and exposure, rate of progression, responses in children, review, subtype comparisons, variant crossrecognition or cross-neutralization

- clone 3: Called CL3 here. Nabs against HIV-1 M group isolates were tested for their ability to neutralize 6 randomly selected HIV-1 O group strains. Clone 3 could neutralize some O group strains. CL3 is specific for a linear epitope containing 2 cysteines that generate a loop that could be important during the fusion of the virus with the target cell. This epitope is represented as GCxGxxxCxT HIV-1 group O isolates. Ferrantelli et al. [2004a] (variant cross-recognition or crossneutralization)
- clone 3: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 -604). Only one of these has any neutralizing potential, clone 3. clone 3 neutralized 3 diverse B clade TCLA strains and 3 primary O group strains. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization, review, subtype comparisons)
- clone 3: Monoclonal antibodies to this epitope have distinct phenotypes-41-7 and 1B8.env were found to be not neutralizing, 2F11 possibly enhancing, and clone 3 beneficial. Enshell-Seijffers et al. [2001] (variant cross-recognition or cross-neutralization)
- clone 3: Inhibits replication of three diverse HIV-1 laboratory strains, as well as an AZT-resistant isolate. Cotropia et al. [1996] (variant cross-recognition or cross-neutralization)
- clone 3: Core binding domain gcsgkLIC lack of serological activity to this region correlates with rapid progression in infants (Broliden et al. [1989]) Cotropia et al. [1992]. Broliden et al. [1989]; Cotropia et al. [1992] (antibody binding site definition and exposure, responses in children, rate of progression)

**No.** 710

MAb ID 4

HXB2 Location gp160 (598–604)

Author Location gp41 (598–609)

Epitope CSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

**Species (Isotype)** mouse (IgG2b)

References Bizub-Bender et al. 1994; Oldstone et al.

• There is another MAb with this ID that reacts with integrase. Bizub-Bender et al. [1994]; Oldstone et al. [1991]

• 4: Stimulated by immunization with the peptide: LGLIWGCS-GKLIC (aa 598-609) - poor cross-reactivity with HIV-2 peptide CAFRQVC – slightly more reactive with longer HIV-2 peptide NSWGCAFRQVC. Oldstone et al. [1991]

No. 711

**MAb ID** 41-6

HXB2 Location gp160 (598-604)

Author Location gp41 (598–609)

Epitope CSGKLIC

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgG2b)

References Oldstone et al. 1991

• 41-6: Stimulated by immunization with the peptide: LGLI-WGCSGKLIC (aa 598-609) – poor cross-reactivity with HIV-2 peptide CAFRQVC - slightly more reactive with LGLIWGCS-GKLIC and HIV-2 form NSWGCAFRQVC - disulfide bond between cysteines required. Oldstone et al. [1991]

No. 712

**MAb ID** 41-7

HXB2 Location gp160 (598–604)

Author Location gp41 (605–611)

**Epitope** CSGKLIC

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

References Enshell-Seijffers et al. 2001; Bugge et al. 1990

- 41-7: Monoclonal antibodies to this epitope have distinct phenotypes-41-7 and 1B8.env were found to be not neutralizing, 2F11 possibly enhancing, and clone 3 beneficial. Enshell-Seijffers et al. [2001]
- 41-7: Sera from 6/6 HIV-1 positive, but no HIV-2 positive individuals, interfered with 41-7 binding – Ab does not neutralize. Bugge et al. [1990]

No. 713

**MAb ID** 68.1

HXB2 Location gp160 (598–604)

Author Location gp41 (598-609)

Epitope CSGKLIC

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) mouse (IgM)

References Oldstone et al. 1991

• 68.1: Stimulated by immunization with the peptide: LGLI-WGCSGKLIC (aa 598-609) - cross-reactive with HIV-2 peptide CAFRQVC - more reactive with longer HIV-1 peptide LGLIWGCSGKLIC and HIV-2 peptide NSWGCAFRQVC. Oldstone et al. [1991]

No. 714

**MAb ID** 68.11

HXB2 Location gp160 (598–604)

Author Location gp41 (598–609)

Epitope CSGKLIC

Neutralizing

Immunogen vaccine

*Vector/Type:* peptide *HIV component:* gp41

Species (Isotype) mouse (IgM)

References Oldstone et al. 1991

• 68.11: Stimulated by immunization with the peptide: LGLI-WGCSGKLIC (aa 598-609) - cross-reactive with HIV-2 peptide CAFRQVC - more reactive with longer HIV-1 peptide Research Contact H. Katinger, Inst. Appl. Microbiol., Vienna, LGLIWGCSGKLIC and HIV-2 peptide NSWGCAFRQVC. Oldstone *et al.* [1991]

No. 715

MAb ID 75

HXB2 Location gp160 (598-604) Author Location gp41 (598–609)

**Epitope** CSGKLIC

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) rat (IgG)

References Oldstone et al. 1991

• 75: Stimulated by immunization with the peptide: LGLIWGC-SGKLIC (aa 598-609) - poor cross-reactivity with HIV-2 peptide CAFRQVC - more reactive with longer HIV-2 peptide NSWGCAFRQVC. Oldstone et al. [1991]

**No.** 716

MAb ID polyclonal

HXB2 Location gp160 (598-604)

**Author Location** gp41 (603–609)

Epitope CSGKLIC

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Enshell-Seifffers et al. 2001

• Monoclonal antibodies to this epitope have distinct phenotypes - 41-7 and 1B8.env were found to be not neutralizing, 2F11 possibly enhancing, and clone 3 beneficial - isolated mimotopepresenting phages corresponding to the immunodominant gp41 epitope CSGKLIC were used to study the diversity of polyclonal responses in 30 HIV+ sera, and all but one of the patients reacted showing distinctive variable polyclonal recognition patterns. Enshell-Seijffers et al. [2001]

No. 717

MAb ID 105-732

HXB2 Location gp160 (599-606)

**Author Location** gp41 (601–608 HAM112, O group)

Epitope KGRLICYT

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: O group

HAM112 HIV component: gp160

**Species (Isotype)** mouse (IgG2b $\kappa$ )

References Scheffel et al. 1999

• 105-732: Overlapping peptides based on group O HAM112 Env were tested for MAb reactivity - MAb 105-732 bound to two overlapping peptides. Scheffel et al. [1999]

No. 718

MAb ID 3D6 (IAM 41-3D6)

HXB2 Location gp160 (599-613)

Author Location gp41 (604–617 BH10)

**Epitope** SGKLICTTAVPWNAS

Neutralizing no

**Immunogen** HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

**Ab Type** gp41 cluster I, immunodominant region

Austria and Viral Testing Systems, Houston,

References Gorny & Zolla-Pazner 2004; Finnegan et al.

2002; Cavacini et al. 1999; Cavacini et al. 1998a; Cavacini et al. 1998b; Kunert et al. 1998; Wisnewski et al. 1996; Stigler et al. 1995; Sattentau et al. 1995; Chen et al. 1994b;

He et al. 1992; Felgenhauer et al. 1990

Keywords antibody binding site definition and exposure, antibody sequence, variable domain, kinetics, review, structure

- 3D6: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- 3D6: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I MAbs 50-69, F240, 240-D,3D6, and 246-D recognize a nonhelical hydrophobic region, positions 598-604, that forms a disulfide loop in the six-helix bundle. Cluster II MAbs 98-6 and 126-6 recognized residues 644-663 of gp41, a portion of the second heptad repeat. These MAbs were found to behave similarly, so 50-69 and 98-6 were used as representatives. Exposure of cluster I and cluster II epitopes required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor. Binding to CD4 exposed hidden cluster I and II epitopes. The MAbs were found to bind to gp120/gp41 complexes, not to gp41 after shedding of gp120, and were localized to at fusing-cell interfaces. Kinetic and binding results indicate that these MAbs are exposed in transitional structures during the fusion process, possibly the prehairpin intermediate prior to co-receptor binding, although other intermediate structures may be involved. They do not bind once syncytia begin to show extensive cyotplasmic mixing. These MAbs failed to inhibit fusion. The NAb 2F5 has a very different behavior in this study. Finnegan et al. [2002] (antibody binding site definition and exposure, kinetics)
- 3D6: Cavacini et al. note that both MAbs F223 and 3D6 are anti-HIV-1 Env MAbs that have an autoimmune response and that both use uses VH3 germline genes. Cavacini et al. [1999]
- 3D6: Binds to the immunodominant region of gp41 a strong homology between heavy variable domains of hu MAb 3D6 and MAb F20 was observed, these MAbs may define a human Ab clonotype. Cavacini et al. [1998a] (antibody sequence, variable domain)
- 3D6: The complete V, J and D(H) domain was sequenced in contrast the sequences of five neutralizing MAbs, 3D6 had very little somatic mutation, with homologies of 97-98% relative

variable domain)

- 3D6: 3D6 is V H3 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996] (antibody sequence, variable domain)
- 3D6: Called IAM 41-3D6: binding increased after pretreatment of infected cells with sCD4 – binding domain overlaps site that is critical for gp120-gp41 association. Sattentau et al. [1995] (antibody binding site definition and exposure)
- 3D6: Optimum peptide for binding 3D6 Fab was CSGKLICT-TAVPW. Stigler et al. [1995] (antibody binding site definition and exposure)
- 3D6: This MAb binds to HIV gp41, and to a 43 kd protein found in human T, B and monocyte cell lines, proposed molecular mimicry. Chen et al. [1994b]
- 3D6: Fab fragment crystal structure. He et al. [1992] (structure)
- 3D6: Sequence of cDNA encoding V-regions. Felgenhauer et al. [1990] (antibody sequence, variable domain)

**No.** 719

MAb ID F172-D8 (F172-D8, scFvD8)

**HXB2 Location** gp160 (604–615)

Author Location gp41 (609–620)

**Epitope** CTTAVPWNASWS?

Neutralizing **Immunogen** 

Species (Isotype) human

References Legastelois & Desgranges 2000

• F172-D8: As an approach to intercellular immunization using a single-chain variable fragment, scFvD8 was constructed based on the MAb F172-D8, directed at a loop in gp41 between the two heptad repeat regions - intracellular scFvD8 expression decreased gp160 expression and a scFvD8 transfected cell line did not support infection by HIV-1 Ba-L or primary isolates. Legastelois & Desgranges [2000]

No. 720

MAb ID D50

HXB2 Location gp160 (632-655)

Author Location gp41 (642–665)

**Epitope** 

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component:

dimeric Env

Species (Isotype) mouse

**Ab Type** gp41 cluster II

Research Contact Patricia Earl and Christopher Broder, NIH

**References** de Rosny et al. 2004a; de Rosny et al. 2004b;

Srivastava et al. 2002; Yang et al. 2000; Earl et al. 1997; Richardson et al. 1996; Binley et al. 1996; Earl et al. 1994

**Keywords** antibody binding site definition and exposure,

antibody generation

- to germline genes. Kunert et al. [1998] (antibody sequence, D50: The MAb 2F5 binds to the C-heptad and is neutralizing, but the MAb D50 binds to the C-heptad and is not neutralizing. 2F5 binds preferentially to native gp41 prior to receptor activation. D50 prefers the triggered form after receptor activation. Trapped fusion-intermediates suggest 2F5 remains present shortly after gp120 triggering by CD4, but may be lost by the time the six-helix bundle is formed. D50 binds equally to the fusion-intermediate and six-helix bundle. 2F5 neutralization seems to block a later step of the fusion process. de Rosny et al. [2004b] (antibody binding site definition and exposure)
  - D50: The mechanism of 2F5 neutralization was explored, and experiments suggest it is due to interference with a late step in viral entry. sCD4 binding to gp120 triggers conformational changes in gp41 allowing formation of the six helix bundle. The NAb 2F5 preferentially bound native gp41, prior to receptor triggering, while the antibody D50 that also binds to the heptad region, near 2F5, is not neutralizing, and preferentially bound the CD4 triggered gp41. The C and N peptides that can be used to block the formation of the six helix bundle and lock gp41 in the fusion intermediate state after sCD4 triggering enabled 2F5 to bind after sCD4 triggering, while D50 was able to bind to both the peptide-trapped and sCD4 induced six helix bundle equally well, suggesting the D50 epitope is linear and more exposed after sCD4 binding. de Rosny et al. [2004a] (antibody binding site definition and exposure)
  - D50: Oligomeric gp140 (o-gp140) derived from R5 primary isolate US4 was characterized for use as a vaccine reagent -D50 was used to capture the o-gp140 for ELISA to test the antigencity of o-gp140 using a panel of well characterized MAbs. Srivastava et al. [2002]
  - D50: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) - gp41 MAbs T4, D12, T3, and D50 bound less efficiently to gp140-GNC4 than did pooled sera, but T4 and D12 recognized the gp140-GNC4 timer equivalently to gp140(-), and T3 and D50 recognized the trimer at greater levels than gp140(-). Yang et al. [2000] (antibody binding site definition and exposure)
  - D50: Found to bind to a linear peptide, between Env amino acids 642-655 - can be blocked by the conformation dependent MAbs D16, D17, D31, D36, D37, D40, D44, D55, D59, T37, and T45 - the region is in the immunogenic cluster two region reactive with 9/10 HIV-1 strains tested, all except HIV-1 ADA, in which the change E659D and E662A may result in the loss of binding (ELLE to DLLA). Earl et al. [1997] (antibody binding site definition and exposure)
  - D50: Thought to be a discontinuous epitope recognizing residues between 649-668 - designated cluster II - Fabs D5, D11, G1, T3, M12, M15, S6, S8, S9, S10 block binding. Binley et al. [1996] (antibody binding site definition and exposure)
  - D50: Richardson suggests this is a linear gp41 epitope. Richardson et al. [1996] (antibody binding site definition and exposure)
  - D50: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994] (antibody generation)

No. 721 MAb ID 5-21-3

**HXB2 Location** gp160 (642–665)

**Author Location** gp41 (642–665 HXB2)

Epitope IHSLIEESQNQQEKNEQELLELDK

Subtype B Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: gp41

Species (Isotype) mouse

References Scheffel et al. 1999; Hunt et al. 1990

- 5-21-3: Binds group M gp41, used as a control in a study of group O MAbs. Scheffel *et al.* [1999]
- 5-21-3: Recognizes a contiguous, conformation-dependent epitope in a hydrophilic region. Hunt *et al.* [1990]

**No.** 722

**MAb ID** 120-16 (SZ-120.16)

HXB2 Location gp160 (644-663)

Author Location gp41 (644–663 HXB2)

Epitope SLIEESQNQQEKNEQELLEL

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human ( $IgG2\kappa$ )

**References** Wisnewski *et al.* 1996; Forthal *et al.* 1995; Eddleston *et al.* 1993; Robinson *et al.* 1991; Xu *et al.* 1991; Tyler *et al.* 1990; Robinson *et al.* 1990b; Andris *et al.* 1992

- 120-16: 120-16 is V H4 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski *et al.* [1996]
- 120-16: No neutralizing activity, both ADCC and viral enhancing activity. Forthal et al. [1995]
- 120-16: Called SZ-120.16. Eddleston et al. [1993]
- 120-16: Synergizes with huMAb 50-69 *in vitro* to enhance HIV-1 infection. Robinson *et al.* [1991]
- 120-16: Less reactive region than AVERY region most Abs involving this region bound conformational epitopes, this was the only linear one. Xu *et al.* [1991]
- 120-16: Antibody dependent enhancement (ADE) of HIV-1 IIIB infectivity, synergistically enhanced by MAb V10-9. Robinson *et al.* [1990b]
- 120-16: Potent ADCC (in contrast to MAb 98-43, gp41(579-604)). Tyler *et al.* [1990]

**No.** 723

**MAb ID** 98-6 (SZ-98.6, 98.6, 98-6D)

HXB2 Location gp160 (644–663)

**Author Location** gp41 (644–663 HXB2)

Epitope SLIEESQNQQEKNEQELLEL

Subtype B

Neutralizing no

**Immunogen** HIV-1 infection **Species (Isotype)** human ( $IgG2\kappa$ )

**Ab Type** gp41 alpha-helical hairpin intermediate, gp41

cluster II

**Research Contact** Susan Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu), NYU, NY

References Ling et al. 2004; Gorny & Zolla-Pazner 2004; Finnegan et al. 2002; Follis et al. 2002; Golding et al. 2000b; Verrier et al. 2001; Taniguchi et al. 2000; Nyambi et al. 2000; Gorny et al. 2000; Gorny & Zolla-Pazner 2000; Nyambi et al. 1998; Hioe et al. 1997b; Wisnewski et al. 1996; Sattentau et al. 1995; Manca et al. 1995a; Forthal et al. 1995; Chen et al. 1995; Laal et al. 1994; Tani et al. 1994; Spear et al. 1993; Eddleston et al. 1993; Xu et al. 1991; Robinson et al. 1991; Sattentau & Moore 1991; Andris et al. 1992; Tyler et al. 1990; Robinson et al. 1990b; Till et al. 1989; Gorny

Keywords ADCC, antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain, binding affinity, complement, enhancing activity, immunotoxin, kinetics, review, subtype comparisons, variant cross-recognition or cross-neutralization

et al. 1989; Pinter et al. 1989

- 98-6: NIH AIDS Research and Reference Reagent Program: 1240.
- 98-6: One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have any neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- 98-6: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of 17b and F105 was decreased by trypsin, but increased by thrombin. gp41 MAbs 246D, 98.6, 50-69, were decreased by trypsin, unaltered by thrombin, while NAb 2F5 binding was increased by thrombin. Thrombin may increase HIV-induced cell fusion in blood by causing a conformational activating shift in gp120. Ling et al. [2004] (antibody binding site definition and exposure)
- 98-6: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I MAbs 50-69, F240, 240-D,3D6, and 246-D recognize a nonhelical hydrophobic region, positions 598-604, that forms a disulfide loop in the six-helix bundle. Cluster II MAbs 98-6 and 126-6 recognized residues 644-663 of gp41, a portion of the second heptad repeat. These MAbs were found to behave similarly, so 50-69 and 98-6 were used as representatives. Exposure of cluster I and cluster II epitopes required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor. Binding to CD4 exposed hidden cluster I and II epitopes. The MAbs were found to bind to gp120/gp41 complexes, not to gp41 after shedding of gp120, and were localized to at fusing-cell interfaces. Kinetic and binding results indicate that these MAbs are exposed in transitional structures during the fusion process, possibly the prehairpin intermediate prior to co-receptor binding, although other intermediate structures may be involved. They do not bind once syncytia begin to show extensive cyotplasmic mixing.

These MAbs failed to inhibit fusion. The NAb 2F5 has a very different behavior in this study. Finnegan *et al.* [2002] (antibody binding site definition and exposure, kinetics)

- 98-6: Called 98-6D. Alanine mutations were introduced into the N- and C-terminal alpha-helices of gp41 to destabilize interhelical packing interactions in order to study their inhibitory effect on viral infectivity. These mutations were shown to inhibit viral replication though affecting the conformational transition to the fusion-active form of gp41, and allow increased inhibition by gp41 peptides. 2F5 senstivity is increased in the mutated viruses, presumably because 2F5s neutralization activity is focused on the transition to the fusion active state. No other gp41 MAb against tested, including NC-1, 50-69D, 1281, 98-6D, 246-D and F240, neutralized the parental or the fusion-deficient mutated viruses. Follis *et al.* [2002] (antibody binding site definition and exposure)
- 98-6: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion 98-6 binds to a C-HR hairpin epitope and blocks fusion when added to a 2 hour E/T preincubation at 31.5 C, but if added after 1 hour, doesn't inhibit this is in contrast to six-helix bundle Abs 167-D and 1281 that inhibit more efficiently when added after one hour of incubation. Golding *et al.* [2002b] (antibody binding site definition and exposure)
- 98-6: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6—six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D, while six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier *et al.* [2001] (antibody interactions, variant cross-recognition or cross-neutralization)
- 98-6: 98-6 and 2F5 both bind to a peptide N51-C43 complex trimer of heterodimers that approximates the core of the fusogenic form of gp41, and to C43 alone but not to N51 alone 98-6 and 2F5 have comparable affinities for C43, but 98-6 has a higher affinity for the complex and the binding of 98-6 is not inhibited by N51. Gorny & Zolla-Pazner [2000] (antibody binding site definition and exposure, binding affinity)
- 98-6: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, but gp41 MAb 50-69 bound with a 5 fold preference for the oligomer, while other gp41 MAbs (1367, 98-6, 167-D, 1281, 1342, and 1379) did not show a preference. Gorny et al. [2000] (antibody binding site definition and exposure)
- 98-6: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 cluster II anti-gp41 MAbs of these 2F5, 167-D, 126-6, and 1281 bound across clades, but usually weakly, while 98-6 and 1342 had poor cross reactivity Clade D isolates bound most consistently to cluster II MAbs no neutralizing activity was observed when tested against 5

isolates, but 98-6 did not bind to these isolates. Nyambi *et al.* [2000] (**subtype comparisons**)

- 98-6: The fusogenic form of gp41 is recognized by 98-6, and the epitope is a conformational epitope formed by the interaction of two regions of gp41 which form an alpha-helical bundle. Taniguchi *et al.* [2000] (antibody binding site definition and exposure)
- 98-6: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H anti-gp41 Abs 98-6, 1367 and 1342 were not able to bind detectably with any of the viruses from any clade. Nyambi *et al.* [1998] (variant cross-recognition or cross-neutralization)
- 98-6: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe *et al.* [1997b] (variant cross-recognition or cross-neutralization)
- 98-6: 98-6 is V H4 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski *et al.* [1996] (antibody sequence, variable domain)
- 98-6: One of several anti-gp41 MAbs that bind to a gp41-maltose binding fusion protein designed to study the leucine zipper domain of gp41, showing that the construct has retained aspects of normal gp41 conformation. Chen *et al.* [1995] (antibody binding site definition and exposure)
- 98-6: No neutralizing activity, positive ADCC activity, and no viral enhancing activity. Forthal *et al.* [1995] (ADCC, enhancing activity)
- 98-6: Virions complexed to gp41 Ab facilitate presentation of p66 RT epitopes to Th cells. Manca *et al.* [1995a]
- 98-6: Preferentially recognizes oligomeric form of gp41 enhanced binding to HIV-1 infected cells at 37 degrees relative to 4 degrees addition of sCD4 enhances binding. Sattentau *et al.* [1995] (antibody binding site definition and exposure)
- 98-6: Epitope described as cluster II, 644-663, conformational does not neutralize IIIB or synergize neutralization by anti-V3 MAb 447-52D or by CD4 BS MAbs. Laal et al. [1994] (antibody binding site definition and exposure, antibody interactions, variant cross-recognition or cross-neutralization)
- 98-6: This MAb was expressed as a surface anti-gp41 monoclonal antibody receptor for gp41 on a CD4-negative B-cell line. Transfected cells could bind HIV Envelope, but could not be infected by HIV-1. When CD4 delivered by retroviral constructs was expressed on these cells, they acquired the ability to replicate HIV-1, and sIg/gp41 specifically enhanced viral replication. Tani et al. [1994]

- 98-6: Called SZ-98.6 binds to a conformational domain within aa 644-663 of gp41, and reacts with astrocytes, as do 167-7 and ND-15G1. Eddleston *et al.* [1993] (antibody binding site definition and exposure)
- 98-6: Did not mediate deposition of complement component C3 on HIV infected cells, binding enhanced by sCD4. Spear *et al.* [1993] (**complement**)
- 98-6: No neutralizing or enhancing activity. Robinson et al. [1991] (enhancing activity)
- 98-6: Two fold increase in binding to gp120 in the presence of bound sCD4. Sattentau & Moore [1991] (antibody binding site definition and exposure)
- 98-6: Appeared to be specific for a conformational or discontinuous epitope. Xu et al. [1991] (antibody binding site definition and exposure)
- 98-6: No neutralizing or enhancing activity for HIV-1 IIIB.
   Robinson et al. [1990b] (enhancing activity)
- 98-6: Serves as target for antibody-dependent cellular cytotoxicity, ADCC. Tyler et al. [1990] (ADCC)
- 98-6: Kills HIV-infected cells when coupled to deglycosylated **Research Contact** Susan ricin A chain. Gorny *et al.* [1989] (**immunotoxin**)
- 98-6: Reacts preferentially with gp160 oligomer, compared to gp41 monomer. Pinter *et al.* [1989] (antibody binding site definition and exposure)
- 98-6: Toxic to HIV-infected T cells (H9) and monocytes (U937) when coupled to deglycosylated A chain of ricin. Till *et al.* [1989] (**immunotoxin**)

No. 724

MAb ID 167-7 (SZ-167.7)

HXB2 Location gp160 (644–663)

Author Location gp41 (644–663)

Epitope SLIEESQNQQEKNEQELLEL

**Neutralizing** 

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG2\lambda$ )

Ab Type gp41 cluster II

References Eddleston et al. 1993; Xu et al. 1991

- 167-7: Called SZ-167.7 binds to a conformational domain within aa 644-663 of gp41, and reacts with astrocytes, as do 98-6 and ND-15G1. Eddleston *et al.* [1993]
- 167-7: Specific for a conformational epitope. Xu et al. [1991]

**No.** 725

**MAb ID** ND-15G1 (ND-15GI)

**HXB2 Location** gp160 (644–663)

**Author Location** gp41 (644–663 HXB2)

Epitope SLIEESQNQQEKNEQELLEL

Subtype B

Neutralizing

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

**Ab Type** gp41 cluster II

References Gorny & Zolla-Pazner 2004; Eddleston et al.

1993

**Keywords** antibody binding site definition and exposure,

review

• ND-15G1: One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)

 ND-15G1: Mapped to the conformational epitope within aa 644-663, and reacts with astrocytes, as do 98-6 and 167-7. Eddleston *et al.* [1993] (antibody binding site definition and exposure)

**No.** 726

**MAb ID** 167-D (167)

HXB2 Location gp160 (644-663)

**Author Location** gp41 (644–663 HXB2)

Epitope SLIEESQNQQEKNEQELLEL

Subtype B

Neutralizing no

**Immunogen** HIV-1 infection **Species (Isotype)** human (IgG1 $\lambda$ )

**Ab Type** gp41 cluster II, gp41six-helix bundle

Research Contact Susan Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu), NYU, NY

References Gorny & Zolla-Pazner 2004; Golding et al. 2002b; Nyambi et al. 2000; Gorny et al. 2000; Gorny & Zolla-Pazner 2000; Manca et al.

1995a; Forthal *et al.* 1995; Spear *et al.* 1993

**Keywords** ADCC, antibody binding site definition and exposure, complement, enhancing activity, review, subtype comparisons, variant cross-recognition or cross-neutralization

- 167-D: Called 167. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- 167-D: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion 98-6 binds to a C-HR hairpin epitope and blocks fusion when added to a 2 hour E/T preincubation at 31.5 C, but if added after 1 hour, doesn't inhibit this is in contrast to six-helix bundle Abs 167-D and 1281 that inhibit more efficiently when added after one hour of incubation. Golding *et al.* [2002b] (antibody binding site definition and exposure)
- 167-D: This cluster II MAb binds to a conformational epitope in the region 644-663 like most cluster II MAbs (126-6, 167-D, 1281, 1342, and 1379 all reacted similarly) it binds to a peptide N51-C43 complex trimer of heterodimers that approximates the core of the fusogenic form of gp41, but not to C43 nor to N51 alone. Gorny & Zolla-Pazner [2000] (antibody binding site definition and exposure)
- 167-D: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, but gp41 MAb 50-69 bound with a 5 fold preference for the oligomer, while other gp41 MAbs (1367, 98-6, 167-D, 1281, 1342, and 1379) did not show a preference. Gorny *et al.* [2000] (antibody binding site definition and exposure)

- 167-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 cluster II anti-gp41 MAbs of these 2F5, 167-D, 126-6, and 1281 bound across clades, but usually weakly, while 98-6 and 1342 had poor cross reactivity Clade D isolates bound most consistently to cluster II MAbs. Nyambi *et al.* [2000] (**subtype comparisons**)
- 167-D: No neutralizing activity, no ADCC activity, and no viral enhancing activity. Forthal *et al.* [1995] (ADCC, enhancing activity, variant cross-recognition or cross-neutralization)
- 167-D: Virions complexed to gp41 Ab facilitate presentation of p66 RT epitopes to Th cells. Manca *et al.* [1995a]
- 167-D: Did not mediate deposition of complement component C3 on HIV infected cells complement mediated virolysis of MN and IIIB in the presence of sCD4. Spear *et al.* [1993] (**complement**)

**No.** 727

MAb ID polyclonal

HXB2 Location gp160 (659-670)

Author Location gp41 (659–670)

**Epitope** ELLELDKWASLW

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: B clade HIV

component: gp41 Adjuvant: QS21

Species (Isotype) guinea pig

References McGaughey et al. 2003

**Keywords** antibody binding site definition and exposure, binding affinity, vaccine antigen design

2F5: Cyclic peptides ELLELDKWASLW that adopt constrained beta-turn conformation of the 2F5 epitope beta-turn in the complexed crystal structure were synthesized and optimize 2F5 binding affinity. This peptide elicits high titer peptide-specific immune responses in guinea pigs that do not neutralize; the authors propose this may be the result of a short CDR3 loop in guinea pigs and additional recessed contact points between 2F5 and gp41. McGaughey et al. [2003] (antibody binding site definition and exposure, vaccine antigen design, binding affinity)

**No.** 728

MAb ID polyclonal

HXB2 Location gp160 (662–667)

Author Location gp41 (662–667)

Epitope ELDKWA

Neutralizing no

Immunogen vaccine

HIV component: gp41

Species (Isotype) guinea pig

References Joyce et al. 2002

• 2F5: DP178 is a peptide derived from the C-term heptad repeat of gp41 that is a potent inhibitor of viral-mediated fusion – it contains ELDKWA but fails to stimulate 2F5-like NAbs upon immunization – the peptide was extended to force an increase in helicity, and the modified peptide had a increase in affinity for 2F5, but upon guinea pig immunization although high peptide-specific Ab titers were achieved the sera were incapable of viral neutralization – the authors propose that 2F5 may be a low affinity maturation intermediate, which may account for its

breadth and why it is hard to recreate the NAb response, but also suggests that the high concentrations required for neutralization are not relevant *in vivo*. Joyce *et al.* [2002]

No. 729

MAb ID polyclonal

HXB2 Location gp160 (662–667)

**Author Location** gp41

Epitope ELDKWA

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41 Adjuvant: Complete Freund's Adjuvant (CFA), Incomplete Freund's Adjuvant (IFA)

Species (Isotype) mouse

References Liu et al. 2005b

**Keywords** vaccine antigen design, vaccine-specific epitope characteristics

A peptide containing eight copies of the MAb 2F5's ELDKWA-epitope separated by as spacers GSGGGGS, RS, and GS was used to test the impact of spacers on eliciting antibody responses to peptides. Both GSGGGGS and GS induced high titers of ELDKWA peptide-specific Abs in BALB/c mice, which reacted with rsgp41. Liu et al. [2005b] (vaccine antigen design, vaccine-specific epitope characteristics)

**No.** 730

MAb ID 5B2

**HXB2** Location gp160 (662–667)

Author Location Env (669–674 IIIB)

Epitope ELDKWA

Neutralizing

Immunogen vaccine

Vector/Type: peptide keyhole limpet hemocyanin (KLH) conjugate Strain: B clade

IIIB HIV component: gp41

**Species (Isotype)** mouse (IgG) **Ab Type** C-domain

References Tian et al. 2001

- 5B2: There is an RT specific Ab Szilvay *et al.* [1992] and a gp41 specific Ab Tian *et al.* [2001] both called 5B2. Tian *et al.* [2001]
- 5B2: Peptides GPGRAFY and ELDKWA were conjugated to keyhole limpet hemocyanin and used to raise mouse MAbs MAb hybridomas were generated with defined specificity 5B2 and 9G11 bind to the peptide and to rgp41. Tian *et al.* [2001]

**No.** 731

MAb ID 9G11

HXB2 Location gp160 (662–667)

**Author Location** Env (669–674 IIIB)

Epitope ELDKWA

Neutralizing

Immunogen vaccine

Vector/Type: peptide keyhole limpet hemocyanin (KLH) conjugate Strain: B clade

IIIB HIV component: gp41

Species (Isotype) mouse (IgG)

Ab Type C-domain References Tian et al. 2001

• 9G11: Peptides GPGRAFY and ELDKWA were conjugated to KLH and used to raise mouse monoclonal Ab-MAb hybridomas were generated with defined specificity—5B2 and 9G11 bind to the peptide and to rgp41. Tian et al. [2001]

No. 732

MAb ID TH-Ab1

HXB2 Location gp160 (662-667)

Author Location gp41 (669-674)

Epitope ELNKWA **Neutralizing** LP

Immunogen vaccine

Vector/Type: peptide keyhole limpet hemocyanin (KLH) conjugate Strain: B clade TH936705 HIV component: gp41 Adjuvant: Complete Freund's Adjuvant (CFA)

Species (Isotype) rabbit (IgG1)

Ab Type C-domain

References Dong et al. 2001; Xiao et al. 2000a

• TH-Ab1: ELNKWA is an escape variant not recognized by the broadly neutralizing MAb 2F5, which recognizes the core epitope ELDKWA-Abs were raised against the peptide escape variant CGELNKWAGELNKWA linked to KLH carrierthese polyclonal antibodies, like the MAb TH-Ab1 also raised to ELNKWA, could recognize ELDKWA and escape mutant peptide epitopes ELEKWA and ELDEWA. Dong et al. [2001]

**No.** 733

MAb ID polyclonal

HXB2 Location gp160 (662–667)

**Author Location** gp41

Epitope ELDKWA

Neutralizing LP

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) rabbit

Ab Type C-domain

References Liao et al. 2000

• Low levels of anti-ELDKWA antibodies are observed in HIV-1 + individuals, so a C-domain P2 peptide linked to a carrier was used to immunize mice and rabbits, and stimulated a high-level anti-ELDKWA response in mice and rabbits - vaccine was C-TSLIHSLIEESQNQQEKNEQELLELDKWA linked to carrier peptide K/G [(KGGG)\_7-K]. Liao et al. [2000]

**No.** 734

MAb ID polyclonal

HXB2 Location gp160 (662-667)

Author Location gp41 (669–674)

Epitope ELDKWA

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: Env

Adjuvant: BSA

Species (Isotype) rabbit, mouse

Ab Type C-domain

References Xiao et al. 2000b

• Strong epitope-specific neutralizing antibody responses were induced using a Env peptide bound to BSA, C(ELDKWAG)\_4-BSA, but not full gp160. Xiao et al. [2000b]

No. 735

MAb ID polyclonal

HXB2 Location gp160 (662–667)

**Author Location** gp41 (662–667 BH10)

Epitope ELDKWA

Neutralizing L

Immunogen vaccine

Vector/Type: influenza Strain: B clade

BH10 HIV component: gp41

Species (Isotype) mouse (IgA, IgG)

Ab Type C-domain

References Muster et al. 1995; Muster et al. 1994

· Sustained ELDKWA specific IgA response in mucosa of immunized mice. Muster et al. [1995]

No. 736

MAb ID polyclonal

HXB2 Location gp160 (662-667)

**Author Location** gp120 (669–674)

Epitope ELDKWA

Neutralizing

Immunogen vaccine

Vector/Type: protein, polyepitope HIV com-

ponent: gp160 Adjuvant: BSA

Species (Isotype) rabbit

Ab Type C-domain

References Lu et al. 2000b; Lu et al. 2000c

 High titer response to ELDKWA and RILAVERYLKD was observed upon vaccination with multiple-epitope vaccine CG-GPGRAFY-G-ELDKWA-G-RILAVERYLKD conjugated to BSA, with a weak response to GPGRAFY - immunization with CG-(ELDKWA-GPGRAFY)\_2-K was also tried, yielding a strong Ab response to both ELDKWA and GPGRAFY – gp160 vaccination yielded strong Ab response but not to any of the peptides studied here. Lu et al. [2000c,b]

No. 737

**MAb ID** 14D9

HXB2 Location gp160 (662–667)

Author Location gp41 (669-674 MVP5180)

Epitope ELDEWA

Subtype B, CRF01\_AE, O

Neutralizing

Immunogen vaccine

Vector/Type: peptide keyhole limpet hemocyanin (KLH) conjugate Strain: natural variants HIV component: gp41 Adjuvant:

Complete Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgG1)

Ab Type gp41 adjacent to cluster II, C-term

References Huang et al. 2002

Keywords antibody binding site definition and expo-

sure, antibody generation, subtype comparisons, variant cross-recognition or cross-

neutralization

• 14D9: This mouse MAb was raised against a variant of ELD-KWA core epitope of the NAb 2F5, eldEwa, derived from the 2F5 neutralization resistance variant MVP5180. The eldEwa peptide was conjugated to the carrier protein keyhole limpet hemocyanin (KLH) and adminstered to BALB/c mice and 14D9 was prepared using standard hybridoma methods. 2F5 does not bind to the variants eldEwa, elNkwa (B.TH.TH936705) or elEkwa, while 14D9 binds only to eldEwa and not ELDKWA. The eldEwa variant is common in the HIV-1 O group. Huang et al. [2002] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 738

**MAb ID** 2F5 (IAM 2F5, IAM-41-2F5, IAM2F5, c2F5)

HXB2 Location gp160 (662-667)

Author Location gp41 (662–667 BH10)

Epitope ELDKWA

Neutralizing LP

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG $3\kappa$ )

Ab Type gp41 adjacent to cluster II, C-term

Research Contact Hermann Katinger, Institute of Applied Microbiology, Vienna, or Polymun Scientific

Inc., Vienna, Austria, or Viral Testi

References Zwick et al. 2005; Wang et al. 2005; Srisurapanon et al. 2005; Ren et al. 2005; Pinter et al. 2005; Nakowitsch et al. 2005; Lusso et al. 2005; Luo et al. 2006; Louis et al. 2005; Liu et al. 2005b; Krachmarov et al. 2005; Ho et al. 2005; Haynes et al. 2005; Chakrabarti et al. 2005; Dong et al. 2005; Biron et al. 2005; Zwick et al. 2004; Menendez et al. 2004; Safrit et al. 2004; Pugach et al. 2004; Pinter et al. 2004; Opalka et al. 2004; Nabatov et al. 2004; McCaffrey et al. 2004; Lorin et al. 2004; Ling et al. 2004; Liao et al. 2004; Jeffs et al. 2004; Ferrantelli et al. 2004a; Ferrantelli et al. 2004b; de Rosny et al. 2004a; de Rosny et al. 2004b; Binley et al. 2004; Gorny & Zolla-Pazner 2004; Wolbank et al. 2003; Ohagen et al. 2003; Montefiori et al. 2003; McGaughey et al. 2003; Mascola et al. 2003; Kitabwalla et al. 2003; Wang 2003; Richman et al. 2003; Mascola 2003; Hart et al. 2003; Ferrantelli et al. 2003; Dey et al. 2003; Binley et al. 2003; Stiegler et al. 2002; Li et al. 2002; Huang et al. 2002; Gorry et al. 2002; Finnegan et al. 2002; Follis et al. 2002; Cavacini et al. 2002; Bures et al. 2002; Liu et al. 2002; Ferrantelli & Ruprecht 2002; Zhang et al. 2002; Kunert et al. 2002; Mascola 2002; Grundner et al. 2002; Xiang et al. 2002b; Clerici et al. 2002a; Joyce et al. 2002; Chakrabarti et al. 2002; Xu et al. 2002; Ho et al. 2002; Tian et al. 2002; Schulke et al. 2002; Golding et al. 2002b; Srivastava et al. 2002; Armbruster et al. 2002;

Root et al. 2001; Xu et al. 2001; Hofmann-Lehmann et al. 2001; Stiegler et al. 2001; Verrier et al. 2001; Spenlehauer et al. 2001; Parker et al. 2001; Zeder-Lutz et al. 2001; Moore et al. 2001; Barnett et al. 2001; Mascola & Nabel 2001; Zwick et al. 2001c; Zwick et al. 2001b; York et al. 2001; Tumanova et al. 2001; Kolchinsky et al. 2001; Dong et al. 2001; Si et al. 2001; Yang et al. 2000; Xiao et al. 2000c; Coeffier et al. 2000; Sanhadji et al. 2000; Pai et al. 2002; Park et al. 2000; Nyambi et al. 2000; Lu et al. 2000b; Lu et al. 2000c; Liao et al. 2000; Kunert et al. 2000; Gorny & Zolla-Pazner 2000; Robert-Guroff 2000; Baba et al. 2000; Mascola et al. 2000; Mascola et al. 1999; Parren et al. 1999; Muhlbacher et al. 1999; Beddows et al. 1999; Poignard et al. 1999; Montefiori & Evans 1999; Frankel et al. 1998; Kunert et al. 1998; Geffin et al. 1998; Parren et al. 1998b; Jiang et al. 1998; Li et al. 1998; Takefman et al. 1998; Ernst et al. 1998; Fouts et al. 1998; Trkola et al. 1998; Yang et al. 1998; Parren et al. 1998a; Connor et al. 1998; Mondor et al. 1998; Andrus et al. 1998; Gorny et al. 1997; Earl et al. 1997; Burton & Montefiori 1997; Ugolini et al. 1997; Turbica et al. 1997; Stamatatos et al. 1997; Mascola et al. 1997; Moore & Trkola 1997; Kessler II et al. 1997; Li et al. 1997; Mo et al. 1997; D'Souza et al. 1997; Schutten et al. 1997; Purtscher et al. 1996; Stoiber et al. 1996; McKeating et al. 1996; Pincus et al. 1996; Conley et al. 1996; Sattentau 1996; Poignard et al. 1996b; McKeating 1996; Calarota et al. 1996; Kessler et al. 1995; Neurath et al. 1995; Moore & Ho 1995; Sattentau et al. 1995; Trkola et al. 1995; D'Souza et al. 1995; Beretta & Dalgleish 1994; Muster et al. 1994; McGaughey et al. 2004; Chen et al. 1994b; Thali et al. 1994; Conley et al. 1994b; D'Souza et al. 1994; Dacheux et al. 2004; Buchacher et al. 1994; Laal et al. 1994; Purtscher et al. 1994; Klasse et al. 1993a; Allaway et al. 1993; Muster et al. 1993; Buchacher et al. 1992

Keywords acute/early infection, adjuvant comparison, antibody binding site definition and exposure, assay standardization/improvement, coreceptor, escape, immunoprophylaxis, immunotherapy, kinetics, mimotopes, motherto-infant transmission, neutralization potency, rate of progression, reversion, viral fitness, review, structure, subtype comparisons, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-neutralization

- 2F5: UK Medical Research Council AIDS reagent: ARP3063.
- 2F5: NIH AIDS Research and Reference Reagent Program: 1475.

- 2F5: gp41 and p15E of the porcine endogenous retrovirus (PERV) share structural and functional similarities, and epitopes in the membrane proximal region of p15E are able to elicit NAbs upon immunization with soluble p15E. Rabbits immunized with a VSV recombinant expressing an HIV-1 membrane-proximal external region (MPER) fused to PERV p15E, with a fusion p15E-HIV MPER protein boost, elicited HIV specific NAbs. The MPER contains the 2F5 epitope, and the 2F5 MAb was used as a positive control for neutralization in this study, and could bind to the vaccine construct. Luo *et al.* [2006] (vaccine antigen design)
- 2F5: Circular dichroism and NMR were used to analyze the structure of the HIV-1 inhibitor peptide T-20 (gp41 HXB2 aa 638-673) that contains the full 2F5 and partial 4E10 epitope. T-20 was unstructured towards the N terminus, and helical in the central and C-terminal regions. The 2F5 epitope sequence (gp41 HXB2 657-670) forms an intrinsic helical structure, which is stable in water. Biron et al. [2005] (structure)
- 2F5: Guinea pigs were immunized with a hybrid HXB2/BaL Env (HIV HXB/BaL gp140δCFI, clade B) in which the tip of the V3 loop (GPGRA) was replaced with the 2F5 epitope LELDKWAS. 2F5 bound to the Env that carried the V3replacement 2F5 epitope, but antibodies against this construct only neutralized the X4-tropic lab adapted HIV strain IIIB, and not CCR5-HIV BaL or SF162 isolates. Chakrabarti et al. [2005] (vaccine antigen design, variant cross-recognition or cross-neutralization)
- 2F5: 2F5 recognizes the epitope ELDKWA, but does not neutralize viruses carrying the commonly found mutated epitope variants: ELDEWA, ELDSWA, ELDNWA, ELDQWA, ELDTWA, or ELNKWA. Peptide cocktails containing ELDKWA, ELNKWA, ELDEWA, and ELEKWA elicit polyclonal antibodies in rabbits that can bind to all of the natural variants that are escape variants for 2F5 expressed in gp41 via WB, as well as ELDERWA. Dong *et al.* [2005] (vaccine antigen design, variant cross-recognition or cross-neutralization, escape)
- 2F5:2F5 and 4E10 both bind to membrane proximal regions of gp41, and have long hydrophobic CDR3 regions characteristic of polyspecific autoreactive antibodies. Of 35 Env-specific MAbs tested, only 2F5 and 4E10 were reactive with phospholipid cardiolipin. Vaccine induction of antibodies that react with these gp41 membrane proximal regions may be rare because of elimination due to autoantigen mimicry. 2F5 also reacted with centromere B and histone autoantigens, and both 4E10 and 2F5 reacted with HEp-2 cells with diffuse cytoplasmic and nuclear patterns indicating polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)
- 2F5: In an attempt to elicit 2F5-like antibodies, the 2F5 epitope ELDKWAS was constrained in the beta-turn sites of the immunoglobulin heavy chain, or alternatively was attached at the C-terminal ends of the immunoglobulin light chain. The constrained heavy chain inserted epitopes bound to 2F5 with 10-fold higher affinity than the light chain unconstrained versions, and when used as an immunogen, elicited epitope-specific antibodies in rabbits, but these antibodies could not neutralize the virus. Ho et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)

- 2F5: Sera from subtype A infected individuals from Cameroon have antibodies that react strongly with subtype A and subtype B V3 loops in fusion proteins, and neutralize SF162 pseudotypes, while sera from 47 subtype B infected individuals reacted only with subtype B V3s. Sera from Cameroon did not neutralize primary A or B isolates, due to indirect masking by the V1/V2 domain rather than due to loss of the target epitope. Neutralization by Cameroonian sera MAbs was blocked by Clade A and B V3 loop fusion proteins, while NAbs to non-V3 epitopes, 2F5, 2G12, and b12, were not blocked. Krachmarov et al. [2005] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, subtype comparisons)
- 2F5: A peptide containing eight copies of the ELDKWA-epitope separated by aa spacers GSGGGGS, RS, and GS was used to test the impact of spacers on eliciting antibody responses to peptides. Both GSGGGGS and GS induced high titers of ELDKWA peptide-specific Abs in BALB/c mice, which reacted with rsgp41. 2F5 served as a positive control in a Western Blot to determine whether epitope-specific Abs bound to recombinant protein rsgp41. Liu et al. [2005b] (vaccine antigen design, vaccine-specific epitope characteristics)
- 2F5: Nine anti-gp41 bivalent Fabs that interacted with either or both of the 6-helix bundle and the internal coiled-coil of N-helices of gp41 were selected from a non-immune human phage display library. The IC50 range for the inhibition of LAV ENV-mediated cell-fusion was 6-61 ug/ml. For context, 2F5 and 2G12 (IC50s of 0.5-1.5 ug/ml) were about an order of magnitude more potent in this assay than the best Fabs generated here. Louis *et al.* [2005] (**neutralization potency**)
- 2F5: Passive immunization of 8 HIV-1 infected patients with 4E10, 2F5 and 2G12 (day 0, 4E10; days 7, 14 and 21 4E10+2G12+2F5; virus isolated on days 0 and 77) resulted in 0/8 patients with virus that escaped all three NAbs. No viruses fully escaped 2F5, although 5/8 developed a more than 2-fold increase in 2F5 IC50 concentrations at day 77. No changes in the 2F5 epitope were observed in the 77 day study period, although 3 patients had unusual 2F5 epitope sequences to start with (not A/ELDKWA but SLNNWN, ALDTWE, or KFDNWA); all viruses were susceptible to 2F5 neutralization, although to varying degrees. In a companion in vitro study, resistance to a single MAb emerged in 3-22 weeks, but triple combination resistance was slower and characterized by decreased viral fitness. In the core of the 2F5 epitope, LDKW, the L and W were completely conserved in the in vitro study, but 9/13 cases had a D->N change, 1/13 a K->N, and 1/13 a K->Q. The lack of resistance to the combination of MAbs in vivo and the reduced fitness of the escape mutants selected in vitro suggests passive immunotherapy may be of value in HIV infection. Nakowitsch et al. [2005] (escape, immunotherapy)
- 2F5: This study is about the V2 MAb C108g, that is type-specific and neutralizes BaL and HXB2. JR-FL is a neutralization resistant strain; modification of JRFL at V2 positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The modification in V2 also increased neutralization sensitivity to V3 MABs 4117c, 2219, 2191, and 447-52D, but only had minor effects on neutralization by CD4BS MAb 5145A, and broadly neutralizing MAbs IgG1b12, 2G12, and 2F5. Pinter et al. [2005] (antibody

## binding site definition and exposure)

- 2F5: The antibody M2 is specific for a peptide flag inserted into the V4 loop of YU-2, a neutralization resistant variant with a short V4 loop. IgG1b12 and 2F5 could neutralize both the WT YU-2 and the modified variant. The high diversity of V4 suggests it does not play a direct role in receptor binding or viral entry, yet M2, specific for the peptide insert tag, can neutralize the modified virus, demonstrating that neutralizing activity doesn't have to block functionality of the virus. Ren et al. [2005] (neutralization potency)
- 2F5: Ab titers to the 2F5 binding peptide ELDKWA were tested by peptide ELISA in sera from Thais infected with CRF01 virus who were asymptomatic versus those who had AIDS, and antibody titers were found to be significantly lower in AIDS patients. The frequency of recognition of this peptide was low overall (15-35%) in CRF01 infections, as well as infections with clades A-G. Srisurapanon *et al.* [2005] (variant cross-recognition or cross-neutralization, subtype comparisons, rate of progression)
- 2F5: A multi-epitope ELDKWA/ELDEWA string in a glutathione S-transferase (GST) backbone elicited Abs in mice and rabbits that could bind to gp41 carrying either the 2F5 susceptible ELDKWA variant, or the ELDEWA escape variant. Vaccinations with only the ELDKWA epitope or the ELDEWA embedded-peptide constructs yielded type specific Abs. Wang et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics, escape)
- 2F5: Alanine scanning mutations of the 21 amino acid region between positions 660-680 showed that only Ala substitutions in the DKW at the core of the epitope reduced binding, positions llelDKWanlwnwfdisnwlw. No single Ala mutation was resistant to both 2F4 and 4E10. Ala substitutions in 12 of the 20 positions enhanced neutralization sensitivity, LLeLd-kwanLWNWFdIsnWLW.2F5 inhibits the neutralization activity of peptide T20. Zwick *et al.* [2005] (antibody binding site definition and exposure, escape)
- 2F5: 93 viruses from different clades were tested for their neutralization cross-reactivity using a panel of HIV antibodies. 2F5 was cross-reactive with A, B, and E subtype viruses, some D, and no C clade viruses. DKW was defined as the core motif, and was found in only 25% of C clade sequences in the database. It was found in C clade viruses in a country specific manner common in Burundi, Brazil and Ethiopia, rare in Botswana, India, and S. Africa. The potency of the neutralizing activity was somewhat context-dependent. DQW is a common D clade variant from Uganda, and all D viruses in this study were Ugandan. Binley et al. [2004] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 2F5: Env sequences were derived from 4 men at primary infection and four years later; the antigenicity in terms of the ability to bind to 2G12, 2F5 and IgG1b12 was determined. 2G12 bound primarily to late clones in 3 of the 4 patients, and to both early and late in the other patient. Neither 2F5 nor IgG1b12 showed a difference in binding affinity to early or late envelopes. Dacheux *et al.* [2004] (antibody binding site definition and exposure, acute/early infection, kinetics)
- 2F5: Neonatal rhesus macaques were exposed orally to a pathogenic SHIV, 89.6P. 4/8 were given an intramuscular, passive immunization consisting of NAbs 2G12, 2F5 and 4E10, each

- given at a different body sites at 40 mg/kg per Ab, at one hour and again at 8 days after exposure to 89.6P. The four animals that were untreated all died with a mean survival time of 5.5 weeks, the four animals that got the NAb combination were protected from infection. This model suggests antibodies may be protective against mother-to-infant transmission of HIV. Ferrantelli *et al.* [2004b] (**mother-to-infant transmission**)
- 2F5: Nabs against HIV-1 M group isolates were tested for their ability to neutralize 6 randomly selected HIV-1 O group strains. IgG1b12 could neutralize some O group strains when used on its own, and quadruple combination of b12, 2F5, 2G12, and 4E10, could neutralize the six Group O viruses tested between 62-97%. The 2F5 epitope in the O group viruses was: ELDEWA. Ferrantelli *et al.* [2004a] (variant cross-recognition or cross-neutralization)
- 2F5: This paper reviews MAbs that bind to HIV-1 Env. 2F5 binds to a region of gp41 proximal to cluster II (aa 662-676), neighboring the binding site of the broadly neutralizing MAb 4E10 and of neutralizing Fab Z13. 2F5 is broadly neutralizing. Gorny & Zolla-Pazner [2004]
- 2F5: A set of oligomeric envelope proteins were made from six primary isolates for potenial use as vaccine antigens: 92/UG/037 (clade A), HAN2/2 (clade B), 92/BR25/025 (clade C), 92/UG/021 (clade D), 93/BR/029 (clade F) and MVP5180 (clade O). This was one of a panel of MAbs used to explore folding and exposure of well characterized epitopes. The clade C isolate BR25 is apparently misfolded, as conformation-dependent antibodies did not bind to it. 2F5 bound to clade A, B, D and F HIV-1 primary isolates. Polyclonal sera raised in rabbits against these antigens cross-bound the other antigens, but none of the sera had neutralizing activity. Jeffs et al. [2004]
- 2F5: 2F5 was used as a positive control in a study that showed that A32-rgp120 complexes open up the CCR5 co-receptor binding site, but did not induce neutralizing antibodies with greater breadth among B subtype isolates than did uncomplexed rgp120 in vaccinated guinea pigs. Liao et al. [2004]
- 2F5: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of 17b and F105 was decreased by trypsin, but increased by thrombin. gp41 MAbs 246D, 98.6, 50-69, were decreased by trypsin, unaltered by thrombin, while NAb 2F5 binding was increased by thrombin. Thrombin may increase HIV-induced cell fusion in blood by causing a conformational activating shift in gp120. Ling et al. [2004] (antibody binding site definition and exposure)
- 2F5: Mice susceptible to MV infection were intraperitoneally immunized with native HIV-1 89.6 env gp160 and gp140 and δV3 HIV-1 89.6 mutants expressed in live attenuated Schwarz measles vector (MV). The gp160ΔV3 construct raised more cross-reactive NAbs to primary isolates. The constructs had an additional 2F5 MAb epitope, ELDKWAS, but responses were not directed towards this epitope. A HIVIG/2F5/2G12 combination was used as a positive control and could neutralize all isolates. Lorin et al. [2004] (vaccine antigen design)

- 2F5: Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The loss of the glycan within the V3 loop (GM299 V3) and adjacent to the C-terminal end of the V3 loop (GM329 C3) did not alter neutralization susceptibility to 2F5, but the loss of glycans in C2 (GM292 C2), C4 (GM438 C4), or V5 (GM454 V5) increased 2F5 neutralization susceptibility. V3 glycans tended to shield V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans shielded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004] (antibody binding site definition and exposure, vaccine antigen design)
- 2F5: This review summarizes properties of 2F5 and its binding to the prefusogenic membrane proximal region of gp41.
   The linear core epitope does not stimulate cross-reactive NAbs when placed outside the context of gp41, suggesting its presentation in a highly specific moleuclar framework is critical.
   McGaughey et al. [2004] (vaccine antigen design, review)
- 2F5: 2F5 was used for screening of phage-displayed peptide libraries. 2F5 requires the DKW core for synthetic and phage-displayed peptide recognition, but is multispecific for amino acid residues flanking C-terminally the DKW core epitope. Three clones from the AADKW-X12 library had high affinity for 2F5, but did not share obvious holomogy with gp41 or each other; Ala substitution showed each bound to 2F5 with a different mechanism. Menendez *et al.* [2004] (antibody binding site definition and exposure, mimotopes)
- 2F5: A set of HIV-1 chimeras that altered V3 net charge and glycosylation patterns in V1V2 and V3, involving inserting V1V2 loops from a late stage primary isolate taken after the R5 to X4 switch, were studied with regard to phenotype, coreceptor usage, and MAb neutralization. The loops were cloned into a HXB2 envelope with a LAI viral backbone. It was observed that the addition of the late-stage isolate V1V2 region and the loss of V3-linked glycosylation site in the context of high positive charge gave an X4 phenotype. R5X4, R5, and X4 viruses were generated, and sCD4, 2G12 and b12 neutralization resistance patterns were modified by addition of the late stage V1V2, glycosylation changes, and charge in concert, while neutralization by 2F5 was unaffected. Nabatov et al. [2004] (antibody binding site definition and exposure, co-receptor)
- 2F5: An antigen panel representing different regions of gp41 was generated, and sera from 23 individuals were screened. Anti-gp41 titers were very high, and sera bound to many regions of gp41, there were no immunologically silent regions. Many individuals had broad responses to diverse regions. High titer responses tended to focus on the N-heptad, C-heptad and 2F5-4E10 regions, but there was no correlation between neutralization capacity of sera and the particular peptides recognized. 2F5 responded to the four antigens that carried the minimal EDLKWA epitope. 2F5 did not bind to the minimal epitope embedded in an alpha helix, supporting that the 2F5 conformation of EDLKWA is embedded in a beta sheet. 2F5 bound better to a synthetic peptide containing the proximal regions

- than to the native gp41. Opalka *et al.* [2004] (assay standardization/improvement)
- 2F5: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Pinter *et al.* [2004] (variant cross-recognition or cross-neutralization)
- 2F5: A primary isolate, CC1/85, was passaged 19 times in PBMC and gradually acquired increased sensitivity to FAb b12 and sCD4 that was attributed to changes in the V1V2 loop region, in particular the loss of a potential glycosylation site. The affinity for sCD4 was unchanged in the monomer, suggesting that the structural impact of the change was manifested at the level of the trimer. The passaged virus, CCcon19, retained an R5 phenotype and its neutralization susceptibility to other Abs was essentially the same as CC1/85. The IC50 for 2F5 was greater than 50 for CC1/85, and was 35 for CCcon19, so the passaged virus was weakly neutralized by 2F5. Pugach et al. [2004] (reversion, viral fitness, variant cross-recognition or cross-neutralization)
- 2F5: The MAb 2F5 binds to the C-heptad and is neutralizing, but the MAb D50 binds to the C-heptad and is not neutralizing. 2F5 binds preferentially to native gp41 prior to receptor activation. D50 prefers the triggered form after receptor activation. Trapped fusion-intermediates suggest 2F5 remains present shortly after gp120 triggering by CD4, but may be lost by the time the six-helix bundle is formed. D50 binds equally to the fusion-intermediate and six-helix bundle. 2F5 neutralization seems to block a later step of the fusion process, but it does not inhibit binding of NC-1, a MAb specific for the six-helix bundle, so it does not prevent formation of the six-helix bundle. The results are most consistent with 2F5 inhibiting a post-fusion-intermediate step. de Rosny et al. [2004b]
- 2F5: The mechanism of 2F5 neutralization was explored, and experiments suggest it is due to interference with a late step in viral entry. sCD4 binding to gp120 triggers conformational changes in gp41 allowing formation of the six helix bundle. The NAb 2F5 preferentially bound native gp41, prior to receptor triggering, while the antibody D50 that also binds to the heptad region, near 2F5, is not neutralizing, and preferentially bound the CD4-triggered gp41. The C and N peptides that can be used to block the formation of the six helix bundle and lock gp41 in the fusion intermediate state after sCD4 triggering enabled 2F5 to bind after sCD4, while D50 was able to bind to both the peptide-trapped and sCD4 induced six helix bundle equally well. The peptide-trapping studies suggest that 2F5 does not fix Env in the native conformation, but interferes with entry after the initial conformation changes occur. Nor does it block six-helix bundle formation, as 2F5 prebinding does not inhibit NC-1 binding, a MAb that binds specifically to the six-helix bundle. de Rosny et al. [2004a]
- 2F5: This review discusses research presented at the Ghent Workshop of prevention of breast milk transmission and immunoprophylaxis for HIV-1 in pediatrics (Seattle, Oct. 2002),

and makes the case for developing passive or active immunoprophylaxis in neonates to prevent mother-to-infant transmission. Macaque studies have shown that passive transfer of NAb combinations (for example, IgG1b12, 2G12, 2F5, and 4E10; or 2G12 and 2F5) can confer partial or complete protection to infant macaques from subsequent oral SHIV challenge. Safrit *et al.* [2004] (**immunoprophylaxis**, **mother-to-infant transmission**)

- 2F5: A complex of the epitope peptide ELDKWAS bound to 2F5 was crystalized, and the peptide was found to interact with amino acids near the base of the very long (22 residue) CDR 3H region of the Ab. Ala substitution of the CDR H3 region confirmed the importance of these sites near the base of the H3 loop for interaction with the epitope in the context of intact gp41 as well as the peptide. A Phe at the apex of the loop was not located directly in the binding site, however binding of 2F5 to the epitope was very sensitive to non-conservative substitutions in this position (F100G, F100H, and F100R); these diminished both binding affinity and 2F5 neutralization, suggesting a role for the very long CDR 3H region. The authors suggest that particularly long CDR H3 regions may be a common feature of HIV-1 NAbs, based on the 22 residues in H3 of 2F5, the 18 H3 residues in b12, and the 22 H3 residues in X5. They express concern that because small animals like mice are unable to elicit Ab responses with such long H3s, they may be poor model systems for HIV vaccine studies. Zwick et al. [2004]
- 2F5: The SOS mutant envelope protein introduces a covalent disulfide bond between gp120 surface and gp41 transmembrane proteins into the R5 isolate JR-FL by adding cysteines at residues 501 and 605. Pseudovirions bearing this protein bind to CD4 and co-receptor bearing cells, but do not fuse until treatment with a reducing agent, and are arrested prior to fusion after CD4 and co-receptor engagement. gp41 NAbs 2F5 and 4E10 are able to potently neutralize the SOS pseudovirion post-attachment, although 2F5 performed relatively poorly in the pre-attachment assay, a further support for previous studies that indicated it does not bind well to native Env, and may bind best after the virus is attached to cells. Binley et al. [2003]
- 2F5: A sCD4-17b single chain chimera was made that can bind to the CD4 binding site, then bind and block co-receptor interaction. This chimeric protein is a very potent neutralizing agent, more potent than IgG1b12, 2G12 or 2F5 against Ba-L infection of CCR5-MAGI cells. It has potential for prophylaxis or therapy. Dey et al. [2003]
- 2F5: Four newborn macaques were challenged with pathogenic SHIV 89.6 and given post exposure prophylaxis using a combination of NAbs 2F5, 2G12, 4E10 and IgG1b12. 2/4 treated animals did not show signs of infection, and 2/4 macaques maintained normal CD4+ T cell counts and had a lower delayed peak viremia compared to the controls. Ferrantelli *et al.* [2003]
- 2F5: This study investigates the effects of glycosylation inhibitors on the binding between HIV-1 gp120 and mannose-binding lectin (MBL). Mannosidase I inhibitor deoxymanno-jirimycin (dMM) inhibits formation of complex and hybrid N-linked saccharides and yields virus with more mannose residues. dMM added during viral production significantly

- enhanced the binding 2F5 and 2G12, but not IgG1b12 in a viral capture assay. Hart *et al.* [2003]
- 2F5: MAbs IgG1b12, 2G12, 2F5 and 4E10 were tested for their ability to neutralize two primary HIV-1 clade A isolates (UG/92/031 and UG/92/037) and two primary HIV-1 clade D isolates (UG/92/001 and UG/92/005). 4E10 demonstrated the most potent cross-neutralization activity. Quadruple administration of MAbs IgG1b12, 2G12, 2F5, and 4E10 induced strong synergistic neutralization of 4 clade A isolates (UG/92/031, UG/92/037, RW/92/020 and RW/92/025) as well as 5 clade D isolates (UG/92/001,UG/9/005, /93/086/RUG/94/108, UG/94/114). The authors note this combination of 4 MAbs neutralizes primary HIV A, B, C, and D isolates. Kitabwalla et al. [2003]
- 2F5: This review dicusses the importance and function of protective antibody responses in animal model studies in the context of effective vaccine development. SHIV models have shown protection using high levels of MAbs can prevent infection, and partial protection that can influence disease course can be obtained from modest levels of NAbs. SHIV challenges studies conducted with infusions of combinations of MAbs b12, 2G12, and 2F5 are reviewed. Mascola [2003]
- 2F5: Infusions of 2F5 and 2G12 intravenously administered 24h prior to vaginal SHIV-89.P challenge are able to protect macaques from infections. Animals that recieve a IL-2 adjuvanted DNA immunization SIV Gag and HIV Env have T-cell responses and lower viral loads, but were not protected. Suboptimal levels of 2F5 and 2G12 were not able to confer sterile protection in combination with the T-cell responses stimulated by DNA immunizations. Mascola et al. [2003] (adjuvant comparison, vaccine-specific epitope characteristics)
- 2F5: Cyclic peptides ELLELDKWASLW that adopt constrained beta-turn conformation of the 2F5 epitope beta-turn in the complexed crystal structure were synthesized and optimize 2F5 binding affinity. This peptide elicits high titer peptide-specific immune responses in guinea pigs that do not neutralize; the authors propose this may be the result of a short CDR3 loop in guinea pigs. McGaughey et al. [2003]
- 2F5: AC10 is a subject who was given treatment early after infection, and had a viral rebound after cessation of therapy, which then declined to a low level. The polyclonal sera from AC10 could potently neutralize the rebound virus, and NAb escape followed with a neutralizing response against the escape variant and subsequent escape from that response. Viral loads remained low in this subject despite escape. The rebound isolate that was potently neutralized by autologous sera was not particularly neutralization sensitive, as it resisted neutralization by sCD4 and MAbs IgG1b12, 2G12 and 2F5, and was only moderately sensitive to sera from other HIV+ individuals that had high titers of NAbs to TCLA strains. Montefiori et al. [2003]
- 2F5: Env genes derived from uncultured brain biopsy samples from four HIV-1 infected patients with late-stage AIDS were compared to env genes from PBMC samples. Brain isolates did not differ in the total number or positions of N-glycosylation sites, patterns of coreceptor usage, or ability to be recognized by gp160 and gp41 MAbs. 2F5 recognized most variants from 3/4 individuals by gp41 WB; the 4th individual had the ELDKWA

variant Aldkwa in all three isolates. The other single Env that was not recognized carried eldRwa. Ohagen *et al.* [2003]

- 2F5: Most plasma samples of patients from early infection had NAb responses to early autologous viruses, and NAbs against heterologous strains tended to be delayed. Serial plasma samples were tested against serial isolates, and neutralization escape was shown to be rapid and continuous throughout infection. Autologous neutralization-susceptible and resistant viruses from four patients were tested for susceptability to neutralizing Ab responses using MAbs 2G12, IgG1b12 and 2F5. No correlation was established, all viruses tested were susceptible to at least one of the neutralizing MAbs. Two patients that did not have an autologous NAb response also did not evolve changes in susceptibility to these MAbs, while one patient with a pattern of autologous neutralization and escape acquired a 2G12 sensitive virus at month 6, and lost IgG1b12 sensitivity at month 21. Richman et al. [2003]
- 2F5: Review of current neutralizing antibody-based HIV vaccine candidates and strategies of vaccine design. Strategies for targeting of the epitopes for NAbs 2F5, 2G12, 4E10, b12, and Z13 are described. Wang [2003]
- 2F5: The broadly neutralizing antibodies 2F5 and 2G12 were class-switched from IgG to IgA and IgM isotypes. Neutralizing potency was increased with valence for 2G12 so the IgM form was most potent, but for 2F5 the IgG form was most potent. Eight primary isolates were tested including two subtype A isolates. The polymeric IgM and IgA Abs, but not the corresponding IgGs, could interfere with HIV-1 entry across a mucosal epithelial layer, although they were limited in a standard neutralization assay. All isotypes could interact with activated human sera, presumably through complement, to inhibit HIV replication. Wolbank et al. [2003]
- 2F5: A combination of MAbs 2F5 and 2G12 given in multiple infusions was found to be safe and well tolerated even in high doses in a phase I study of seven HIV-1 infected healthy volunteers—the median elimination half-life was 7.94 days for 2F5, and 16.48 for 2G12—no anti-2F5 or anti-2G12 IgM or IgG responses were detected—although there was some transient increases, overall plasma viral RNA levels decreased in 6/7 volunteers, by a median of 0.62 log\_10. Armbruster *et al.* [2002]
- 2F5: IgG1b12 neutralized many South African (5/8) and Malawian (4/8) clade C primary HIV-1 isolates, being more effective than 2F5 which neutralized only two Malawian and no South African isolates. 2G12 did not neutralize any of the 16 isolates. Bures *et al.* [2002]
- 2F5: This study examined antibody interactions, binding and neutralization with a B clade R5 isolate (92US660) and R5X4 isolate (92HT593). Abs generally bound and neutralized the R5X4 isolate better than the R5 isolate, with the exception of F240 which bound both equally well, which captured more virus than any other human MAb tested, and didn't neutralize either isolate. F240 enhanced the binding of CD4BS MAbs IgG1b12 and F105 and the gp41 MAb 2F5 for both R5X4 and R5 isolates. F240 also enhanced neutralization of the R5X4 isolate by 2F5, but had no effect on R5 virus. Anti-V3 MAb B4a1 did not impact 2F5 neutralization. Cavacini *et al.* [2002]
- 2F5: A modified gp140 (gp140deltaCFI), with C-term mutations intended to mimic a fusion intermediate and stabilize

- trimer formation, retained antigenic conformational determinants as defined by binding to CD4 and to MAbs 2F5, 2G12, F105, and b12, and enhanced humoral immunity without diminishing the CTL response in mice injected with a DNA vaccine. Chakrabarti *et al.* [2002]
- 2F5: Six sera from HIV-exposed uninfected individuals(EU) had IgA neutralizing activity dominated by recognition of a distinctive epitope within gp41, QARILAV sera of QAFILAV-immunized BALB/c mice was neutralizing with the dose-dependent behavior similar to 2F5. Clerici et al. [2002a]
- 2F5: Review of NAbs that notes that 2F5 alone or in combination with other MAbs can protect some macaques against SHIV infection, that it is safe and well tolerated in humans, and that illustrates gp41's conformational change and exposure of the 2F5 epitope in the transient pre-hairpin form. Ferrantelli & Ruprecht [2002]
- 2F5: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I and Cluster II MAbs required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor, binding to a fusion intermediate. 2F5 behaved very differently than these non-neutralizing antibodies: it bound to Env in the absence of target cells, and it was distributed evenly all over the cell surface, not localized in fusion domains. It did not interact with cells that exhibited cytoplasmic mixing. 2F5 was unusual in that it exhibited temperature dependence, and did not interact below 19 degrees C, in contrast to 2G12, M77 98-6 and IgG1b12 which bound strongly at temperatures ranging between 4-37 degrees. The authors suggest the temperature dependence of 2F5 may be due to encreased flexibility of the Envelope spike at warmer temperatures facilitating epitope exposure. Finnegan et al. [2002]
- 2F5: Alanine mutations were introduced into the N- and C-terminal alpha-helices of gp41 to destabilize interhelical packing interactions in order to study their inhibitory effect on viral infectivity. These mutations were shown to inhibit viral replication though affecting the conformational transition to the fusion-active form of gp41, and allow increased inhibition by gp41 peptides. 2F5 senstivity is increased in the mutated viruses, presumably because 2F5s neutralization activity is focused on the transition to the fusion active state. No other MAb against gp41 tested, including NC-1, 50-69D, 1281, 98-6D, 246-D and F240, neutralized the parental or the fusion-deficient mutated viruses. Follis et al. [2002]
- 2F5: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion the preincubation 31.5 C step did not alter the inhibitory activity of neutralizing Abs anti-gp41 2F5, or anti-gp120 2G12, IG1b12, 48d, and 17b. Golding *et al.* [2002b]
- 2F5: UK1-br and MACS2-br are R5 isolates derived from brain tissue samples from AIDS patients with dementia and HIV-1 encephalitis; both are neurotropic, but only UK1-br induced neuronal apoptosis and high levels of syncytium formation in macrophages. UK1-br Env had a greater affinity for CCR5 than MACS-br, and required low levels of CCR5 and

CD4 for cell-to-cell fusion and single round infection. PBMC infected with UK1-br and MACS2-br virus isolates were resistant to neutralization by MAb 2G12. UK1-br was more sensitive than MACS2-br to IgG1b12, 2F5 and CD4-IgG2 neutralization. This pattern of Ab reactivity was similar to the to CD4-independent varient ADA197N/K, and thought to result from conformational changes which better expose the CCR5 binding regions, although the loss of the particular N-linked glycosylation site in the V1V2 stem region of ADA was experimentally shown to not be responsible for the the CD4-independent phenotype of UK1-br. Gorry *et al.* [2002]

- 2F5: HIV-1 gp160ΔCT (cytoplasmic tail-deleted) proteoliposomes (PLs) containing native, trimeric envelope glycoproteins from R5 strains YU2 and JRFL, and X4 strain HXBc2, were made in a physiologic membrane setting as candidate immunogens for HIV vaccines—2F5 bound to gp160ΔCT with a reconstituted membrane ten-fold better than the same protein on beads (except for the YU2 form that doesn't bind 2F5)—anti-CD4BS MAbs IgG1b12 and F105, A32 (C1-C4), C11 (C1-C5), and 39F (V3) MAbs bound gp160ΔCT PLs indistinguishably from gp160ΔCT expressed on the cell surface. Grundner et al. [2002]
- 2F5: ELDKWAS was embedded into a beta-turn-like conformational site on a framework of an antibody specific for human leukocyte antigen HLA-DR this construct was recognized by 2F5, and is suggested as an adjuvant-independent vaccine candidate. Ho et al. [2002]
- 2F5: A mouse MAb was raised against a variant of ELDKWA core epitope of the NAb 2F5, eldEwa, derived from the 2F5 neutralization resistant variant MVP5180. 2F5 does not bind to the variants eldEwa, elNkwa (B.TH.TH936705) or elEkwa, while 14D9 binds only to eldEwa and not ELDKWA. The eldEwa variant is common in the HIV-1 O group. Huang et al. [2002]
- 2F5: DP178 is a peptide derived from the C-term heptad repeat of gp41 that is a potent inhibitor of viral-mediated fusion—it contains the 2F5 epitope but fails to stimulate 2F5-like NAbs upon immunization—the peptide was extended to force an increase in helicity, and the modified peptide had a increase in affinity for 2F5, but upon guinea pig immunization although high peptide-specific Ab titers were achieved the sera were incapable of viral neutralization—the authors propose that 2F5 may bind with low affinity to a maturation intermediate, which may account for its breadth and why it is hard to recreate the epitope, but also suggests that the high concentrations required for neutralization are not relevant *in vivo*. Joyce *et al.* [2002]
- 2F5: A 2F5 anti-idiotype murine MAb Ab2/3H6 was developed that blocks 2F5 binding to a synthetic epitope peptide and to gp160 in an ELISA competition assay – Ab2/3H6 diminished the neutralizing potency of 2F5 – Ab2/3H6 Fab fragments were capable of inducing neutralizing Abs and 2F5-epitope specific responses in immunized B6D2F1 mice. Kunert *et al.* [2002]
- 2F5: A polyepitope vaccine was designed based on three repeats of the 2F5 core epitope ELDKWA combined with the V3 region peptide GPGRAFY. Abs raised in mice could recognize the peptides, sgp41, and CHO-WT cells that expressed HIV-1 Env on their surface. Li et al. [2002]
- 2F5: Review of NAbs that discusses mechanisms of neutralization, passive transfer of NAbs and protection in animal studies,

and vaccine strategies. Liu et al. [2002]

- 2F5: Rhesus macaques were better protected from vaginal challenge with SHIV89.6D (MAb 2G12, 2/4; MAbs 2F5/2G12, 2/5; and HIVIG/2F5/2G12, 4/5 infected) than from intravenous challenge (MAb 2G12, 0/3; MAbs 2F5/2G12, 1/3; and HIVIG/2F5/2G12, 3/6 infected)— the animals that were infected by vaginal challenge after Ab infusion had low or undetectable viral RNA levels and modest CD4 T-cell decline. Mascola [2002]
- 2F5: ELDKWAS co-crystallized bound to the Fab' 2F5 fragment showed the epitope peptide in a type I beta-turn conformation. Pai et al. [2002]
- 2F5: Ab binding characteristics of SOS gp140 were tested using SPR and RIPA SOS gp140 is gp120-gp41 bound by a disulfide bond NAbs 2G12, 2F5, IgG1b12, CD4 inducible 17b, and 19b bound to SOS gp140 better than uncleaved gp140 (gp140unc) and gp120 non-neutralizing MAbs 2.2B (binds to gp41 in gp140unc) and 23A (binds gp120) did not bind SOS gp140 SOS gp140-2F5-IgG1b12 formed multiple ring structures composed of two SOS gp140 proteins bridged by two Ab molecules, while 2F5 and 2G12 formed extended chains rather than closed rings. Schulke *et al.* [2002]
- 2F5: Oligomeric gp140 (o-gp140) derived from R5 primary isolate US4 was characterized for use as a vaccine reagent antigen capture ELISA was used to compare the antigenicity of gp120 and o-gp140 using a panel of well characterized MAbs 2F5 recognized o-gp140. Srivastava et al. [2002]
- 2F5: The antiviral response to intravenously administered MAbs 2F5 and 2G12 was evaluated in 7 HAART-naive asymptomatic HIV-1 infected patients during a treatment period of 28 days. MAb therapy reduced plasma HIV RNA in 3/7 patients during the treatment period, and transiently reduced viral load in two more. CD4 counts were up in 3/7 through day 28, and transiently increased in three more. Vigorous complement activation was observed after 48/56 Ab infusions. Before treatment, 2F5 neutralized isolates from five patients and no escape was observed during treatment. Stiegler et al. [2002]
- 2F5: Expanding the minimal epitope ELDKWA to an end-capped, linear nonapeptide, Ac-LELDKWASL-amide attained maximal affinity within a set of native gp41-sequence peptides scanning single residue substitutions confirmed that essential recognition requirements were the central DKW core sequence and the importance of the terminal Leu residues for high-affinity binding high specificity binding pockets at central Lys and Trp side-chains and an absolute requirement for the carboxylate group of the Asp side chain were found the nine residue fragment flanked by pairs of Ser and constrained by a disulfide bridge had high affinity for 2F5. Tian et al. [2002]
- 2F5: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations 375 S/W seems to favor a conformation of gp120 closer to the CD4-bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5,

or CD4i antibodies, but did bind to CD4BS MAbs. Xiang *et al.* [2002b]

- 2F5: Passive immunization of neonate macaques with a combination of F105+2G12+2F5 conferred complete protection against oral challenge with SHIV-vpu+ or the combination b12+2G12+2F5 conferred partial protection against SHIV89.6 such combinations may be useful for prophylaxis at birth and against milk born transmission the synergistic combination of IgG1b12, 2G12, 2F5, and 4E10 neutralized a collection of HIV clade C primary isolates. Xu et al. [2002]
- 2F5: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002]
- 2F5: ELNKWA is an escape variant not recognized by the broadly neutralizing MAb 2F5, which recognizes the core epitope ELDKWA – Abs were raised against the peptide escape variant CGELNKWAGELNKWA linked to KLH carrier – these polyclonal antibodies, like the monoclonal antibody TH-Ab1 also raised to ELNKWA, could recognize ELDKWA and escape mutant peptide epitopes ELEKWA and ELDEWA. Dong et al. [2001]
- 2F5: A combination of MAbs IgG1b12, 2F5, and 2G12 was given postnatally to four neonates macaques that were then challenged with highly pathogenic SHIV89.6P one of the four infants remained uninfected after oral challenge, two infants had no or a delayed CD4(+) T-cell decline. Hofmann-Lehmann et al. [2001]
- 2F5: Mutations in two glcosylation sites in the V2 region of HIV-1 ADA at positions 190 and 197 (187 DNTSYRLINCNTS 199) cause the virus to become CD4-independent and able to enter cells through CCR5 alone – these same mutations tended to increase the neutralization sensitivity of the virus, including to antibody 2F5. Kolchinsky et al. [2001]
- 2F5: Review of studies in macaques that have shown immune control of pathogenic SHIV viremia, improved clinical outcome, and protection, and the implications of the observations for HIV vaccines. Mascola & Nabel [2001]
- 2F5: Moore and colleagues review the data concerning the lack of a clear relationship between genetic subtype and serotype 2F5 is considered in some detail, as it represents a rare vulnerability from the neutralizing antibody perspective, although while it is apparently linear, attempts to present the peptide to the immune system have failed to elicit neutralizing Abs. Moore *et al.* [2001]
- 2F5: Matrix-assisted laser desorption ionization mass spectrometry (MALDI-MS) in combination with proteolytic protection was used to identify the functional epitope for MAb 2F5, NEQELLELDKWASLWN, in the disulfide bond associated gp120/gp41 protein SOS-gp140 (JRFL) this minimal epitope is much larger than the ELDKWA core epitope previously

- defined by peptide ELISA, and this could help explain why ELDKWA-peptides are poor immunogens in terms of eliciting a 2F5-like antibody response. Parker *et al.* [2001]
- 2F5: A peptide called 5-Helix was designed that binds to the C-peptide region of gp41 5-Helix is a potent inhibitor of HIV-1 entry that binds immediately COOH-terminal to the C-peptide region targeted by 5-Helix the conformation of the bound 2F5 epitope is a hairpin turn. Root et al. [2001]
- 2F5: SHIV-HXBc2 is a neutralization sensitive non-pathogenic virus, and several *in vivo* passages through monkey's yielded highly pathogenic SHIV KU-1 HXBc2 and the KU-1 clone HXBc2P3.2 differ in 12 amino acids in gp160 substitutions in both gp120 and gp41 reduced the ability of sCD4, IgG1b12, F105 and AG1121 to Env achieve saturation and full occupancy, and neutralize KU-1 17b and 2F5 also bound less efficiently to HXBc2P3.2, although 2G12 was able to bind both comparably. Si *et al.* [2001]
- 2F5: A luciferase-reporter gene-expressing T-cell line was developed to facilitate neutralization and drug-sensitivity assays luciferase and p24 antigen neutralization titer end points were found comparable using NAb from sera from HIV+ donors, and MAbs 2F5, 2G12 and IgG1b12. Spenlehauer et al. [2001]
- 2F5: 4E10 binds proximal to 2F5 and neutralizes primary isolates of clades A, B, C, D, and E – viruses that were resistant to 2F5 were neutralized by 4E10 and vice versa. Stiegler *et al.* [2001]
- 2F5: A phage peptide library was screened with MAb 2F5, and from the peptides that bound the amino acids DKW were found to be most critical for binding – the mimetic peptide RDWSFDRWSLSEFWL elicited a cross-reactive Ab response to gp41 when used to immunize rabbits. Tumanova et al. [2001]
- 2F5: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001]
- 2F5: Twenty HIV clade C isolates from five different countries were susceptible to neutralization by anti-clade B MAbs in a synergistic quadruple combination of mAbs IgG1b12, 2G12, 2F5, and 4E10. Xu *et al.* [2001]
- 2F5: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding. York et al. [2001]
- 2F5: Neutralizing synergy between MAbs 1b12, 2G12 and 2F5 was studied using surface plasmon resonance to determine the binding kinetics for these three mAbs with respect to monomeric and oligomeric env protein gp160 IIIB – the 2G12 epitope is highly accessible on both monomeric and oligomeric Envs, 1b12 is highly accessible on monomers but not

oligomers, and 2F5 on neither form – binding of 2G12 exposes the 2F5 epitope on gp160 oligomers. Zeder-Lutz *et al.* [2001]

- 2F5: This paper primarily concerns 4E10 and Z13, MAbs that both bind proximally to the 2F5 binding site to a conserved epitope, and that neutralize some primary isolates from clades B, C, and E the minimal 2F5 epitope is determined to be EQELLELDKWASLW, based on screening a gp160 fragment expression library, longer than previous studies broadly neutralizing MAbs 2F5, IgG1b12, and 4E10 and Z13 fail to neutralize different subsets of viruses. Zwick et al. [2001b]
- 2F5: Neutralization synergy between anti-HIV NAbs b12, 2G12, 2F5, and 4E10 was studied a classic fixed-ratio method was used, as well as a method where one Ab was fixed at a low neutralization titer and the other was varied using primary isolates, a two-four fold enhancement of neutralization was observed with MAb pairs, and a ten-fold enhancement with a quadruple Ab combination no synergy was observed with any MAb pair in the neutralization of TCLA strain HXB2. Zwick et al. [2001c]
- 2F5: Paper uses IgG1 form of 2F5 a triple combination of 2F5, F105 and 2G12 effectively neutralized perinatal infection of macaque infants when challenged with SHIV-vpu+ the plasma half-life was 4.2 +/- 0.8 days. Baba *et al.* [2000]
- 2F5: MAbs 98-6 and 2F5 both bind to a peptide N51-C43 complex trimer of heterodimers that approximates the core of the fusogenic form of gp41, and to C43 alone but not to N51 alone 98-6 and 2F5 have comparable affinities for C43, but 98-6 has a higher affinity for the complex and 2F5 may bind to an epitope of C43 that is directly involved with complex formation –and IgG1 rec form of the Ab was used in this study. Gorny & Zolla-Pazner [2000]
- 2F5: 2F5 is a candidate for immunotherapy, but generally IgG1 has a longer half life in humans than IgG3, so the isotype was switched rec CHO-derived MAb 2F5 IgG1kappa and hybridoma-derived MAb 2F5 IgG3kappa displayed identical specificity, *in vitro* function, and epitope (ELDKWA) it remains to be determined if isotype switching will prolongs beta-clearance. Kunert *et al.* [2000]
- 2F5: Low levels of anti-ELDKWA antibodies are observed in HIV-1 + individuals, so a C-domain P2 peptide linked to a carrier was used to immunize mice and rabbits, and stimulated a high-level anti-ELDKWA response. Liao *et al.* [2000]
- 2F5: ELDKWA peptide vaccine study. Lu et al. [2000c]
- 2F5: ELDKWA peptide vaccine study. Lu et al. [2000b]
- 2F5: Because HIV-1 is most often transmitted across mucosal surfaces, the ability of passive transfer of infused HIVIG/2F5/2G12 to protect against mucosal exposure of macaques to pathogenic SHIV 89.6PD was studied HIVIG/2F5/2G12 protected 4/5 animals against vaginal challenge, 2F5/2G12 combined protected 2/5 animals, and 2G12 alone protected 2/4 animals in contrast, Mascola and coworkers had previously shown single MAbs could not protect against intervenous challenge Ab treated animals that got infected through vaginal innoculation had low viral loads and only modest declines in CD4 counts the infused Abs were detected in the nasal, vaginal, and oral mucosa. Mascola et al. [2000]
- 2F5: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 cluster II anti-gp41 MAbs –

- of these 2F5, 167-D, 126-6, and 1281 bound across clades, but usually weakly, while 98-6 and 1342 had poor cross reactivity Clade D isolates bound most consistently to cluster II MAbs. Nyambi *et al.* [2000]
- 2F5: A mini-review of observations of passive administration of IgG NAbs conferring protection against intervenous or vaginal SHIV challenge, that considers why IgG MAbs might protect against mucosal challenge. Robert-Guroff [2000]
- 2F5: 2F5 or sCD4-IgG chimeric immunoadhesin were transferred into 3T3 cells, incorporated into a collagen structure called the neo-organ, and transplanted into SCIDhu mice that were then challenged with MN or LAI the continuous production of the therapeutic molecules in this context resulted in dramatic reduction of viral load. Sanhadji et al. [2000]
- 2F5: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) gp41 MAbs T4, D12, T3, and D50 bound less efficiently to gp140-GNC4 than did pooled sera, but T4 and D12 recognized the gp140-GNC4 timer equivalently to gp140(-), and T3 and D50 recognized the trimer at greater levels than gp140(-) 2F5 did not bind efficiently to these constructs, presumably because of the YU2 strain has a substitution in the 2F5 epitope (ALDKWA instead of ELDKWA). Yang *et al.* [2000]
- 2F5: rgp120 derived from a R5X4 subtype B virus was used to vaccinate healthy volunteers and the resulting sera were compared with sera from HIV-1 positive subjects and neutralizing MAbs. Beddows et al. [1999]
- 2F5: Combinations of HIVIG, 2F5, 2G12 were administered in passive-transfer experiments 24 hours prior to challenge with pathogenic SHIV 89.6PD 3/6 animals given HIVIG/2F5/2G12 were completely protected, the others had reduced viremia and normal CD4 counts 1/3 monkeys given 2F5/2G12 showed transient infection, the other two had reduced viral load all monkeys that received HIVIG, 2F5, or 2G12 alone became infected and developed high-level plasma viremia, although animals that got HIVIG or 2G12 had a less profound CD4 T cell decline. Mascola *et al.* [1999]
- 2F5: A meeting summary presented results regarding neutralization –MAbs 2G12 and 2F5 tested for their ability to neutralize primary isolate infection of genetically engineered cell lines (cMAGI and others, presented by T. Matthews, A. Trkola, J. Bradac) an advantage of such cells lines over PBMCs is that markers (X-Gal) can be added for staining to simplify the assay the consensus of the meeting was that these engineered cell lines did not improve the sensitivity of detection of primary isolate neutralization D. Burton and J. Mascola presented results concerning passive immunization and protection of hu-PBL-SCID mice and macaques, respectively, and both found combinations of MAbs that were able to achieve 99% neutralization *in vitro* corresponded to efficacy *in vivo*. Montefiori & Evans [1999]
- 2F5: In a study of 116 HIV-1 + individuals, Ab reactivity to a peptide encompassing the ELDKWA peptide decreased in CDC stage C patients compared with stage A patients, and longitudinal studies showed a decline in 6/8 patients, while overall Ab reactivity to rec soluble gp160 stayed constant. Muhlbacher et al. [1999]

- 2F5: Review of the neutralizing Ab response to HIV-1. Parren *et al.* [1999]
- 2F5: Hu-PBL-SCID mice were infected with HIV-1s JRCSF and SF162 to study the effect of NAbs on an established infection no significant differences in the initial rate of decrease in viral load or the plateau levels of viral RNA between the b12 treated and control mice were seen in most of the Ab treated mice b12 escape mutants were observed with varying patterns of mutations a combination of b12, 2G12 and 2F5 protected 1/3 mice, and an isolate from one of the other two was resistant to neutralization by all three MAbs. Poignard *et al.* [1999]
- 2F5: Post-exposure prophylaxis was effective when MAb 694/98-D was delivered 15 min post-exposure to HIV-1 LAI in hu-PBL-SCID mice, but declined to 50% if delivered 60 min post-exposure, and similar time constraints have been observed for HIVIG, 2F5 and 2G12, in contrast to MAb BAT123 that could protect delivered 4 hours post infection. Andrus *et al.* [1998]
- 2F5: Ab from gp120 vaccinated individuals prior to infection, who subsequently became HIV infected, could not achieve 90% neutralization of the primary virus by which the individuals were ultimately infected these viruses were not particularly refractive to neutralization, as determined by their susceptibility to neutralization by MAbs 2G12, IgG1b12, 2F5 and 447-52D. Connor et al. [1998]
- 2F5: The ELDKWA epitope was inserted into the antigenic site B of influenza hemagglutinin and expressed on baculovirus infected insect cells, flanked by 3 additional random amino acids, xELDKWAxx – FACS was used to isolate the clone that displayed the epitope with the most markedly increased binding capacity for 2F5, to identify particularly specific immunogenic constructs – PELDKWAPP was a high affinity form selected by FACS. Ernst et al. [1998]
- 2F5: Points out that 2G12 and 2F5, potent neutralizing antibodies, were identified by screening for cell surface (oligomeric Envelope) reactivity. Fouts et al. [1998]
- 2F5: Infection of dendritic cells cultured from CD14+ blood cells or from cadaveric human skin was blocked by neutralizing MAbs IgG1b12, or 2F5 and 2G12 delivered together, but not by control non-neutralizing anti-gp120 MAb 4.8D, indicating that NAbs could interrupt early mucosal transmission events. Frankel et al. [1998]
- 2F5: The natural immune response to the epitope of 2F5, ELD-KWA, was studied in perinatally infected children and levels of reactivity to this epitope were correlated with absolute CD4 numbers over time and health status 3/10 children who had no antibody reactivity to ELDKWA had substitutions in the epitope (ALDKWA, ELDQWA, and KLDKWA) 2F5 competed with the ELDKWA-reactive sera depending on the serum titer. Geffin *et al.* [1998]
- 2F5: Used as a control in the study of anti-gp41 MAb NC-1
   2F5 does not react with HIV-2 gp41 or gp160. Jiang et al.
   [1998]
- 2F5: The complete V, J and D(H) domain was sequenced unlike non-neutralizing anti-gp41 MAb 3D6, five neutralizing MAbs (2F5, 2G12, 1B1, 1F7, and 3D5) showed extensive somatic mutations giving evidence of persistent antigenic pressure over long periods in contrast to Geffin98, where multiple pediatric sera were found to compete with 2F5, cross-

- competition was noted to be very rare in sera from HIV+ adults Kunert *et al.* propose that because there is a binding site of human complement factor H which overlaps the 2F5 binding site, it may generally be masked from the immune system 2F5 also has a remarkably long CDR3 loop of 22 amino acids, and this region could not be readily assigned to any described D(H) fragment, leading to the suggestion of recombination of two fragments from novel regions. Kunert *et al.* [1998]
- 2F5: Neutralization synergy was observed when the MAbs 694/98-D (V3), 2F5 (gp41), and 2G12 (gp120 discontinuous) were used in combination, and even greater neutralizing potential was seen with the addition of a fourth MAb, F105 (CD4 BS). Li et al. [1998]
- 2F5: This MAb and the results of Ugolini *et al.* [1997] are discussed the authors propose that an Ab bound to gp41 would typically project less from the surface of the virion and so be unable to interfere with attachment Parren *et al.* [1998a]. Parren *et al.* [1998a]; Ugolini *et al.* [1997]
- 2F5: MAbs 2G12, 2F5 and b12 are broadly neutralizing, as are some human polyconal sera, but this paper describes a set of primary isolates that are resistant to all three MAbs and 2 broadly neutralizing sera results indicate that resistance levels of pediatric isolates might be higher than adult isolates resistance in general did not seem to be conferred by a loss of binding affinity for gp120 or gp41, rather by a more global perturbation of oligomeric Envelope. Parren et al. [1998b]
- 2F5: Induces complement-mediated lysis in MN but not primary isolates primary isolates are refractive to CML. Takefman et al. [1998]
- 2F5: A wide range of neutralizing titers was observed that was independent of co-receptor usage – 2F5 was the most potent of the MAbs tested. Trkola *et al.* [1998]
- 2F5: A neutralization assay was developed based on heminested PCR amplification of the LTR (HNPCR) LTR-HNPCR consistently revealed HIV DNA and was shown to be a rapid, specific and reliable neutralization assay based on tests with 6 MAbs and 5 isolates. Yang et al. [1998]
- 2F5: This review summarizes results about 2F5: it binds extracellularly, near the transmembrane domain, it is the only gp41 MAb that is neutralizing, it reacts with many non-B clade viruses and has a paradoxically weak binding to virus, given the neutralizing titers. Burton & Montefiori [1997]
- 2F5: In a multilab evaluation of monoclonal antibodies, only IgG1b12, 2G12, and 2F5 could neutralize at least half of the 9 primary test isolates at a concentration of < 25 mug per ml for 90% viral inhibition the isolates with no 2F5 neutralizing susceptibility had the sequences ALGQWA or ELDTWA instead of EDLKWA 7/9 primary isolates were neutralized, and ALDKWQ and ALDKWA were susceptible to neutralization. D'Souza *et al.* [1997]
- 2F5: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB Env strong neutralizer of SHIV-vpu+ all Ab combinations tested showed synergistic neutralization 2F5 has synergistic response with MAbs 694/98-D (anti-V3), 2G12, b12, and F105. Li *et al.* [1997]
- 2F5: Using concentrations of Abs achievable *in vivo*, the triple combination of 2F5, 2G12 and HIVIG was found to be syner-

gistic to have the greatest breadth and magnitude of response against 15 clade B primary isolates. Mascola *et al.* [1997]

- 2F5: A JRCSF variant that was selected for IgG1b12 resistance remained sensitive to MAbs 2G12 and 2F5, for combination therapy. Mo *et al.* [1997]
- 2F5: Review: MAbs 2F5, 2G12 and IgG1b12 have potential for use in combination with CD4-IgG2 as an immunotherapeutic or immunoprophylactic – homologous MAbs to these are rare in humans and vaccine strategies should consider including constructs that may enhance exposure of these MAbs' epitopes. Moore & Trkola [1997]
- 2F5: Called IAM 2F5 antibody mediated enhancement or inhibition seemed to be determined by isolate rather than antibody specificity in this study, only 2F5 inhibited the entry of all the viruses studied, irrespective of their phenotype, and directly proportional to its affinity to monomeric HIV-1 gp160. Schutten *et al.* [1997]
- 2F5: Of three neutralizing MAbs (257-D, IgG1b12, and 2F5),
   2F5 was the only one to inhibit the entry of all viruses studied,
   both SI and NSI, with a potency proportional to its affinity for monomeric gp126. Schutten *et al.* [1997]
- 2F5: Binding of anti-gp120 MAbs IgG1b12 or 654-30D does not mediate significant exposure of the gp41 epitopes for MAbs 2F5 and 50-69. Stamatatos et al. [1997]
- 2F5: Used to standardize polyclonal response to CD4 BS. Turbica *et al.* [1997]
- 2F5: The only MAb out of a large panel to show no correlation between viral binding inhibition and neutralization. Ugolini *et al.* [1997]
- 2F5: IgG1b12 was more potent with greater breadth than MAb 2F5 in an infection reduction assay including 35 primary isolates. Kessler II et al. [1997]
- 2F5: Only 4/20 Argentinian and 3/43 Swedish HIV+ sera reacted with LLELDKWASL sera reacting with peptides that contained ELDKWA tended to have high neutralization titers the region carboxyl terminal to EDLKWA was found to be more important for polyclonal sera AB binding, 670-675 WNWFDI 2F5 bound most strongly to the peptide QELLELDKWA. Calarota *et al.* [1996]
- 2F5: 2F5 was infused into two chimpanzees which were then given an intravenous challenge with a primary HIV-1 isolate

   both became infected, but with delayed detection and prolonged decrease in viral load relative to controls, indicating that preexisting, neutralizing antibodies (passively administered or actively elicited) affect the course of acute-phase virus replication and can be influential after the Ab can no longer be detected in the peripheral circulation. Conley et al. [1996]
- 2F5: Neutralizes HXB2, primary isolates, and chimeric virus with gp120 from primary isolates in an HXB2 background. McKeating et al. [1996]
- 2F5: A panel of immunotoxins were generated by linking Env MAbs to ricin A – immunotoxins mediated cell killing, but killing was not directly proportional to binding. Pincus *et al.* [1996]
- 2F5: Review: one of three MAbs (IgG1b12, 2G12, and 2F5) generally accepted as having significant potency against primary isolates. Poignard *et al.* [1996b]
- 2F5: Primary isolates from clade A, B, and E are neutralized by
   2F5 neutralization requires the LDKW motif neutralization

- resistant isolates or 2F5 selected variants all had substitutions in the D or K. Purtscher *et al.* [1996]
- 2F5: Review: only four epitopes have been described which can stimulate a useful neutralizing response to a broad spectrum of primary isolates, represented by the binding sites of MAbs: 447-52-D, 2G12, Fab b12, and 2F5. Sattentau [1996]
- 2F5: ELDKWAS is in a gp41 binding region for the negative regulator of complement factor H (CFH) Abs to HIV generally do not cause efficient complement-mediated lysis, but binding of 2F5 can interfere with CHF binding, facilitating HIV destruction by complement. Stoiber *et al.* [1996]
- 2F5: Found to neutralize MN, JRCSF, and two B subtype primary isolates, but not a D subtype primary isolate, by most labs in a multi-laboratory study involving 11 labs. D'Souza *et al.* [1995]
- 2F5: Broad cross-clade neutralization of primary isolates additive neutralization in combination with anti-CD4BS MAb IgG1b12 (Called BM12). Kessler *et al.* [1995]
- 2F5: Review: binds to the only generally accepted strong neutralizing epitope outside of gp120, one of only 3 MAbs with strong broad activity against primary viruses, the others are 2G12 and IgG1b12 – unique member of epitope cluster Moore & Ho [1995] and John Moore, per comm 1996. Moore & Ho [1995]
- 2F5: MAb binding decreases the accessibility or alters the conformation of the gp41 fusion domain and of gp120 domains, including the binding site for the CD4 cell receptor. Neurath et al. [1995]
- 2F5: Called IAM 41-2F5 exposed in the presence of gp120 on the cell surface, while most of gp41 is masked binds proximal to transmembrane region. Sattentau *et al.* [1995]
- 2F5: Cross-clade primary virus neutralizing activity LDKW defined as the core epitope. Trkola et al. [1995]
- 2F5: MAb generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994]
- 2F5: Called IAM-41-2F5 neutralized lab and primary isolates

   t 1/2 dissociation 122 min for the peptide, and 156 min for gp41 core D(K/R)W Ab resistant isolate had the sequence KLDNWA. Conley et al. [1994b]
- 2F5: Included in a multi-lab study for antibody characterization binding and neutralization assay comparison. D'Souza *et al.* [1994]
- 2F5: Failed to show synergy with anti-CD4 binding site IIIB neutralizing antibodies. Laal et al. [1994]
- 2F5: 2F5 epitope ELDKWA inserted into an immunogenic loop in influenza virus hemagglutinin can elicit IIIB, MN and RF neutralizing sera in immunized mice. Muster et al. [1994]
- 2F5: Broadly reactive neutralizing activity, ELDKWA is relatively conserved neutralized 2 primary isolates. Purtscher *et al.* [1994]
- 2F5: gp41 mutation (582 A/T) that reduces neutralization of anti-CD4 binding site MAbs does not alter 2F5's ability to neutralize. Thali et al. [1994]
- 2F5: Synergy with combinations of CD4-based molecules in inhibition of HIV-1 Env mediated cell fusion. Allaway *et al*. [1993]
- 2F5: Called IAM-41-2F5 reports MAb to be IgG1 the gp41 mutation 582(Ala to Thr) results in conformational changes in gp120 that confer neutralization resistance to conformationally

sensitive neutralizing MAbs – neutralization efficiency of 2F5 is not affected. Klasse et al. [1993a]

• 2F5: DKWA defined as the core sequence – highly conserved epitope neutralizing MAb. Buchacher et al. [1992]; Muster et al. [1993]

**No.** 739

MAb ID 4E10

HXB2 Location gp160 (671-676)

Author Location gp160 (671-676 MN)

**Epitope NWFDIT** 

Neutralizing P

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG3\kappa$ )

Ab Type C-term

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tria, or Polymum Scientific Inc.,

References Luo et al. 2006; Zwick et al. 2005; Raviv et al. 2005; Nakowitsch et al. 2005; Haynes et al. 2005; Cardoso et al. 2005; Safrit et al. 2004; Pugach et al. 2004; Opalka et al. 2004; Ferrantelli et al. 2004a; Ferrantelli et al. 2004b; Binley et al. 2004; Gorny & Zolla-Pazner 2004; Kitabwalla et al. 2003; Wang 2003; Fiebig et al. 2003; Ferrantelli et al. 2003; Binley et al. 2003; Ferrantelli & Ruprecht 2002; Xu et al. 2002; Xu et al. 2001; Zwick et al. 2001c; Zwick et al. 2001b; Stiegler et al. 2001; D'Souza et al. 1994; Buchacher et al. 1994; Buchacher et al. 1992

Keywords antibody binding site definition and exposure, antibody generation, antibody interactions, assay development, assay standardization/improvement, escape, immunoprophylaxis, immunotherapy, mother-to-infant transmission, reversion, viral fitness, review, structure, subtype comparisons, vaccine antigen design, variant cross-recognition or crossneutralization

- 4E10: gp41 and p15E of the porcine endogenous retrovirus (PERV) share structural and functional similarities, and epitopes in the membrane proximal region of p15E are able to elicit NAbs upon immunization with soluble p15E. Rabbits immunized with a VSV recombinant expressing an HIV-1 membrane-proximal external region (MPER) fused to PERV p15E, with a fusion p15E-HIV MPER protein boost, elicited HIV specific NAbs. The MPER contains the 4E10 epitope. Luo et al. [2006] (vaccine antigen design)
- 4E10: The crystal structure of 4E10 complexed with a 13 aa peptide (KGWNWFDITNWGK) that contains the NWFDIT binding site was resolved to 2.2 A resolution. 4E10 has a canonical beta sandwich Ig-fold, with H3/H2 loop hydophobicity and a long CDR H3 loop that mediates C-terminal base and central amino acid interactions; it extends beyond the peptide and its orientation suggests it could potentially allow hydrophobic contacts with the viral membrane. 4E10 complex formation induces a conformational change in the peptide such that it forms an amphipathic alpha-helix with a hydrophobic face that interacts with 4E10, with Trp672 primary, and Phe673, Ile675

and Thr676 secondary, contact points. Cardoso et al. [2005] (structure)

- 4E10: 2F5 and 4E10 both bind to membrane proximal regions of gp41, and have long hydrophobic CDR3 regions characteristic of polyspecific autoreactive antibodies. Of 35 Env-specific MAbs tested, only 2F5 and 4E10 were found to be reactive with phospholipid cardiolipin. Vaccine induction of antibodies that react with these gp41 membrane proximal regions may be rare because of elimination due to autoantigen mimicry. 4E10 also reacted with systemic lupus erythematosis (SLE) autoantigen SS-A/Ro, and both 4E10 and 2F5 reacted with HEp-2 cells with diffuse cytoplasmic and nuclear patterns indicating polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)
- 4E10: Passive immunization of 8 HIV-1 infected patients with 4E10, 2F5 and 2G12 (day 0, 4E10; days 7, 14 and 21 4E10+2G12+2F5; virus isolated on days 0 and 77) resulted in 0/8 patients with virus that escaped all three NAbs. No viruses escaped 4E10, but only one virus in one patient had the NWFDIT epitope sequence; the W, F and I were conserved in all patients but the other amino acids varied both before and after treatment. A patient carrying the epitope sequence nwfSit had the least 4E10 sensitive virus. In a companion in vitro study, resistance to a single MAb emerged in 3-22 weeks, but triple combination resistance was slower and characterized by decreased viral fitness. In the core of the 4E10 epitope, NWFDIT, 5/11 cases had a T->I escape; 2/11 had a F->L change; and 2/11 had substantial deletions, of WNWF overlapping, or NWLWYI adjacent to the epitope. The lack of resistance to the combination of MAbs in vivo and the reduced fitness of the escape mutants selected in vitro suggests passive immunotherapy may be of value in HIV infection. Nakowitsch et al. [2005] (escape, immunotherapy)
- 4E10: Retrovirus inactivation for vaccine antigen delivery was explored through lipid modification by hydrophobic photoinduced alkylating probe 1.5 iodonaphthylazide (INA). The viral proteins were shown to be structurally intact in the treated noninfectious virus, through the preservation of antibody binding sites for polyclonal anti-gp120 serum, and for broadly neutralizing MAbs 2G12, b12 and 4E10, although the modifications of the lipid disabled viral infection. Raviv et al. [2005] (vaccine antigen design)
- 4E10: Alanine scanning mutations of the 21 amino acid region between positions 660-680 showed only 3 substitutions that reduced 4E10 binding, positions lleldkwanlwnWFdisnwlW. No single Ala mutation was resistant to both 2F4 and 4E10. Ala substitutions in 11/20 positions enhanced neutralization sensitivity, LLeLdkWanLWNwfdIsNWLw. For peptides T20 and 4E10 neutralization was synergistic. Zwick et al. [2005] (antibody binding site definition and exposure, escape)
- 4E10: 93 viruses from different clades were tested for their neutralization cross-reactivity using a panel of HIV antibodies. 4E10 was the most cross-reactive, moderately reactive in all 93 viruses tested from each subtype. WFXI was defined as the core motif, and this core is highly conserved in all M group gp41 sequences. How potent the neutralizing activity is is somewhat context dependent. Binley et al. [2004] (variant cross-recognition or cross-neutralization, subtype comparisons)

- 4E10; Neonatal rhesus macaques were exposed orally to a pathogenic SHIV, 89.6P. 4/8 were given an intramuscular, passive immunization consisting of NAbs 2G12, 2F5 and 4E10, each given at a different body sites at 40 mg/kg per Ab, at one hour and again at 8 days after exposure to 89.6P. The four animals that were untreated all died with a mean survival time of 5.5 weeks, the four animals that got the NAb combination were protected from infection. This model suggests antibodies may be protective against mother-to-infant transmission of HIV. Ferrantelli et al. [2004b] (mother-to-infant transmission)
- 4E10: Nabs against HIV-1 M group isolates were tested for their ability to neutralize 6 randomly selected HIV-1 O group strains. IgG1b12 could neutralize some O group strains when used on its own, and quadruple combination of b12, 2F5, 2G12, and 4E10, could neutralize the six Group O viruses tested between 62-97%. The linear epitope, NWFDIT, of 4E10 is conserved in 3/6 group O strains. Ferrantelli *et al.* [2004a] (variant cross-recognition or cross-neutralization)
- 4E10: This paper reviews MAbs that bind to HIV-1 Env. 4E10 binds to a region of gp41 proximal to cluster II (aa 662-676), neighboring the binding site of the broadly neutralizing MAb 2F5 and overlapping the epitope of neutralizing Fab Z13. 4E10 is the most broadly neutralizing MAb, neutralizing primary isolates from clades A, B, C, D, and CRF01 (E), although not the most potent. Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, subtype comparisons)
- 4E10: An antigen panel representing different regions of gp41 was generated, and sera from 23 individuals were screened. Anti-gp41 titers were very high, and sera bound to many regions of gp41, there were no immunologically silent regions. Many individuals had broad responses to diverse regions. High titer responses tended to focus on the N-heptad, C-heptad and 2F5-4E10 regions, but there was no correlation between neutralization capacity of sera and the particular peptides recognized. 4E10 responded to the three antigens that carried the minimal NWFNIT epitope, but was conformation and context sensitive. Opalka et al. [2004] (assay development, assay standardization/improvement)
- 4E10: A primary isolate, CC1/85, was passaged 19 times in PBMC and gradually acquired increased sensitivity to FAb b12 and sCD4 that was attributed to changes in the V1V2 loop region, in particular the loss of a potential glycosylation site. The affinity for sCD4 was unchanged in the monomer, suggesting that the structural impact of the change was manifested at the level of the trimer. The passaged virus, CCcon19, retained an R5 phenotype and its neutralization susceptibility to other Abs was essentially the same as CC1/85. The IC50 for 4E10 was greater than 50 for CCcon19, and was 44 for CC1/85, so the primary virus was weakly neutralized by 4E10. Pugach et al. [2004] (reversion, viral fitness, variant cross-recognition or cross-neutralization)
- 4E10: This review discusses research presented at the Ghent Workshop of prevention of breast milk transmission and immunoprophylaxis for HIV-1 in pediatrics (Seattle, Oct. 2002), and makes the case for developing passive or active immunoprophylaxis in neonates to prevent mother-to-infant transmission. Macaque studies have shown that passive transfer of NAb combinations (for example, IgG1b12, 2G12, 2F5, and 4E10)

- can confer partial or complete protection to infant macaques from subsequent oral SHIV challenge. Safrit *et al.* [2004] (immunoprophylaxis, mother-to-infant transmission)
- 4E10: The SOS mutant envelope protein introduces a covalent disulfide bond between gp120 surface and gp41 transmembrane proteins into the R5 isolate JR-FL by adding cysteines at residues 501 and 605. Pseudovirions bearing this protein bind to CD4 and co-receptor bearing cells, but do not fuse until treatment with a reducing agent, and are arrested prior to fusion after CD4 and co-receptor engagement. gp41 NAbs 2F5 and 4E10 are able to potently neutralize the SOS pseudovirion post-attachment. Binley et al. [2003] (vaccine antigen design)
- 4E10: Four newborn macaques were challenged with pathogenic SHIV 89.6 and given post exposure prophylaxis using a combination of NAbs 2F5, 2G12, 4E10 and IgG1b12.
   2/4 treated animals did not show signs of infection, and 2/4 macaques maintained normal CD4+ T cell counts and had a lower delayed peak viremia compared to the controls. Ferrantelli *et al.* [2003] (antibody interactions, immunoprophylaxis, mother-to-infant transmission)
- 4E10: Porcine endogenous retroviruses (PERVS) are a concern in the context of porcine xenotransplantation into humans; possible strategies for protection include PERV knockout animals or vaccines. Goats immunized with the PERV transmembrane protein revealed two NAb epitope, E1 and E2. E2's epitope (FEGWFN) binds to a sequence that is perfectly preserved in all PERVS and highly conserved in all gammaretroviruses: MuLV carries FEGLFN, FeLV FEGWFN, and it shares three amino acids with the core epitope for the anti-HIV human neutralizing MAb 4E10, (LWNWFN). Fiebig et al. [2003]
- 4E10: MAbs IgG1b12, 2G12, 2F5 and 4E10 were tested for their ability to neutralize two primary HIV-1 clade A isolates (UG/92/031 and UG/92/037) and two primary HIV-1 clade D isolates (UG/92/001 and UG/92/005). 4E10 demonstrated the most potent cross-neutralization activity. Quadruple administration of MAbs IgG1b12, 2G12, 2F5, and 4E10 induced strong synergistic neutralization of 4 clade A isolates (UG/92/031, UG/92/037, RW/92/020 and RW/92/025) as well as 5 clade D isolates (UG/92/001, UG/9/005, /93/086/RUG/94/108, UG/94/114). The authors note this combination of 4 MAbs neutralizes primary HIV A, B, C, and D isolates. Kitabwalla et al. [2003] (antibody interactions, immunoprophylaxis, variant cross-recognition or cross-neutralization, mother-to-infant transmission, subtype comparisons)
- 4E10: Review of current neutralizing antibody-based HIV vaccine candidates and strategies of vaccine design. Strategies for targeting of the epitopes for NAbs 2F5, 2G12, 4E10, b12, and Z13 are described. Wang [2003] (vaccine antigen design, review)
- 4E10: Review of NAbs illustrating gp41's conformational change and exposure of the 4E10/Z13 epitope in the transient pre-hairpin form. Ferrantelli & Ruprecht [2002] (antibody binding site definition and exposure)
- 4E10: Passive immunization of neonate macaques with a
  combination of F105+2G12+2F5 conferred complete protection against oral challenge with SHIV-vpu+ —the combination b12+2G12+2F5 conferred partial protection against
  SHIV89.6—such combinations may be useful for prophylaxis
  at birth and against milk born transmission—the synergistic

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combination of IgG1b12, 2G12, 2F5, and 4E10 neutralized a collection of HIV clade C primary isolates. Xu et al. [2002] (antibody interactions, immunoprophylaxis, subtype comparisons)

- 4E10: 4E10 binds proximal to 2F5 and neutralizes primary isolates of clades A, B, C, D, and E. Viruses that were resistant to 2F5 were neutralized by 4E10 and vice versa. Stiegler et al. [2001] (antibody binding site definition and exposure)
- 4E10: Twenty HIV clade C isolates from five different countries were susceptible to neutralization by anti-clade B MAbs in a synergistic quadruple combination of mAbs IgG1b12, 2G12, 2F5, and 4E10. Xu et al. [2001] (antibody interactions, subtype comparisons)
- 4E10: MAbs 4E10 and Z13 both bind proximally to 2F5 to a conserved linear epitope that has some conformational aspects - both bind to MN virions, bind weakly to infected cells in a manner that is not disrupted by sCD4 and neutralize some primary isolates from clades B, C, and E – maps minimal 4E10 epitope to NWFDIT, contrary to an earlier report - different strains were refractive to neutralization by broadly neutralizing Abs IgG1b12, 2F5, Z13 and 4E10. Zwick et al. [2001b] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 4E10: Neutralization synergy between anti-HIV NAbs b12, 2G12, 2F5, and 4E10 was studied – a classic fixed-ratio method was used, as well as a method where one Ab was fixed at a low neutralization titer and the other was varied - using primary isolates, a two-four fold enhancement of neutralization was observed with MAb pairs, and a ten-fold enhancement with a quadruple Ab combination - no synergy was observed with any MAb pair in the neutralization of TCLA strain HXB2. Zwick et al. [2001c] (antibody interactions)
- 4E10: MAbs generated by hybridoma, electrofusion of PBL from HIV-1 + volunteers with CB-F7 heteromyeloma cells also binds to MHC class II proteins - anti-class II Abs are only found in HIV-1 positive people - this paper maps 4E10's binding site to AEGTDRV, gp160(823-829), but the later Zwick et al. study in 2001 revised the epitope location. Buchacher et al. [1994] (antibody binding site definition and exposure, antibody generation)
- 4E10: Included in a multi-lab study for antibody characterization, binding and neutralization assay comparison. D'Souza et al. [1994] (variant cross-recognition or crossneutralization)

**No.** 740 MAb ID Z13 HXB2 Location gp160 (671-676) Author Location gp41 (671–676 MN) Epitope NWFDIT Subtype B Neutralizing P Immunogen HIV-1 infection **Species (Isotype)** human (IgG1 $\kappa$ ) **Ab Type** C-term

References Luo et al. 2006; Gorny & Zolla-Pazner 2004;

Zwick et al. 2001b

Keywords antibody binding site definition and exposure, antibody generation, review, vaccine antigen design, variant cross-recognition or cross-neutralization

- Z13:gp41 and p15E of the porcine endogenous retrovirus (PERV) share structural and functional similarities, and epitopes in the membrane proximal region of p15E are able to elicit NAbs upon immunization with soluble p15E. Rabbits immunized with a VSV recombinant expressing an HIV-1 membrane-proximal external region (MPER) fused to PERV p15E, with a fusion p15E-HIV MPER protein boost, elicited HIV specific NAbs. The MPER contains the Z13 epitope. Luo et al. [2006] (vaccine antigen design)
- Z13: This paper reviews MAbs and Fabs that bind to HIV-1 Env. Z13 binds to a region of gp41 proximal to cluster II (aa 662-676), neighboring the binding site of the broadly neutralizing MAb 2F5 and overlapping the epitope of neutralizing MAb 4E10. Z13 is broadly neutralizing, neutralizing primary isolates from clades A, B, C, D and CRF01 (E). Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, review)
- Z13: Review of current neutralizing antibody-based HIV vaccine candidates and strategies of vaccine design. Strategies for targeting of the epitopes for NAbs 2F5, 2G12, 4E10, b12, and Z13 are described. Wang [2003] (vaccine antigen design,
- Z13: Review of NAbs that notes Z13 is a phage display generated FAb fragment from a B clade infected individual and that illustrates gp41's conformational change and exposure of the 4E10/Z13 epitope in the transient pre-hairpin form. Ferrantelli & Ruprecht [2002] (antibody binding site definition and exposure, antibody generation)
- Z13: MAb 4E10 and FAb Z13 both bind proximally to 2F5 to a relatively conserved linear epitope that has some conformational aspects - both bind to MN virions, bind weakly to infected cells in a manner that is not disrupted by sCD4 and can neutralize some primary isolates from clades B, C, and E – Z13 was selected using a phage display library with the MN gp41 peptide LLELDKWASLWNWFDITNWSW from an HIV infected donor who had an exceptionally broad NAb response – different strains were refractive to neutralization by broadly neutralizing Abs IgG1b12, 2F5, Z13 and 4E10 - epitope location noted here is by analogy to MAb 4E10. Zwick et al. [2001b] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization)

No. 741 MAb ID B30 HXB2 Location gp160 (720–734) Author Location gp41 (720–734 BH10) Epitope HLPIPRGPDRPEGIE **Neutralizing** Immunogen vaccine

> Vector/Type: protein Strain: B clade LAI HIV component: gp160

Wang 2003; Ferrantelli & Ruprecht 2002; Species (Isotype) mouse (IgG1) Research Contact George Lewis References Abacioglu et al. 1994

cioglu et al. [1994]

**No.** 742

MAb ID polyclonal

HXB2 Location gp160 (724-745)

Author Location gp41 (731–752)

Epitope PRGPDRPEGIEEEGGERDRDRS

Neutralizing

Immunogen vaccine

Vector/Type: Cowpea mosaic virus Strain:

B clade IIIB HIV component: gp41

Species (Isotype) mouse (IgA, IgG2a)

References Durrani et al. 1998

· Comparison of intranasal and oral immunization of HIV-1 peptide expressed in a plant viral vector - intranasal gave the better response. Durrani et al. [1998]

No. 743

**MAb ID** 41S-2

HXB2 Location gp160 (725-745)

**Author Location** gp160 (732–750)

Epitope RGPDRPEGIEEEGGERDRDRS

**Neutralizing** yes

Immunogen vaccine

Vector/Type: peptide keyhole limpet hemo-

cyanin (KLH) conjugate HIV component:

gp41

**Species (Isotype)** mouse (IgG2b $\kappa$ )

References Hifumi et al. 2003; Hifumi et al. 2002; Hifumi

et al. 2000b; Hifumi et al. 2000a

**Keywords** anti-idiotype, antibody sequence, variable do-

- 41S-2: A murine Ab called i41SL1-2 was raised against the complementary determining region of the 41S-2 light chain, CRDL-1 (RSSKSLLYSNGNTYLY). As with 41S-2-L, the light chain of i41SL1-2 also had catalytic activity and degraded the immunizing peptide, initially cleaving between the Arg1 and Ser2. i41SL1-2 did not cross-react with gp41 peptide, gp120 V3 loop peptide and bound weakly to 41S-2-L.i41SL1-2 shows homology to the anti-VIP Ab (VIP, vasoactive intestinal peptide) that also has peptidase character. Both light chains contain a catalytic triad composed of Asp, Ser, and His (for i41SL1-2: Asp73, Ser 76 or Ser70 and His 79). Intact i41SL1-2 was unable to degrade CDRL-1, possibly due to an immobile inactive conformation of the catalytic triad. Hifumi et al. [2003] (anti-idiotype, antibody sequence, variable domain)
- 41S-2: 41S-2-L refers to the light chain of 41S-2, which can enzymatically decompose the gp41 protein of HIV-1, but doesn't degrade unrealted proteins. The peptide RGPDRPEGIEEEG-GERDRDRS, against which the MAb was raised, can also be cleaved, initially between Glu12-Gly13, followed by successive cleavage reactions. Hifumi et al. [2002]
- 41S-2: BALBc mice were immunized with gp41 peptide and a MAb specific for the peptide was generated – isolated MAb light chains displayed proteolytic activity toward the peptide epitope which may be due to a catalytic triad on light chain (Asp73, Ser76, and His79) – no catalytic activity was observed for the whole antibody. Hifumi et al. [2000a]

• B30: Epitope boundaries mapped by peptide scanning. Aba- • 41S-2: The complementary determining region of 41S-2-L, the light chain of 41S-2, is strongly involved in gp41 recognition. This light chain can serve as a molecular catalyst for gp41 degredation. Hifumi et al. [2000b]

No. 744

MAb ID 447-52D (447/52-DII, 447-52-D, 447d, 447-

52-D, 447-D, 447, 447D)

HXB2 Location gp160 (726-729)

Author Location gp120 (MN)

**Epitope** GPXR

Subtype B

**Neutralizing** LP

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG3 $\lambda$ )

Ab Type gp120 V3

Research Contact Dr. Susan Zolla-Pazner, NYU Med Center NY, NY; Veteran Affairs Med Center NY, NY;

or Cellular Products Inc, Buffalo, NY,

References Selvarajah et al. 2005; Pinter et al. 2005;

Martín-García et al. 2005; Lusso et al. 2005; Krachmarov et al. 2005; Haynes et al. 2005; Gorny et al. 2005; Sharpe et al. 2004; Pugach et al. 2004; Pinter et al. 2004; Pantophlet et al. 2004; McCaffrey et al. 2004; Ling et al. 2004; Gorny et al. 2004; Binley et al. 2004; Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Kessler et al. 2003; Binley et al. 2003; Poignard et al. 2003; Fer-

rantelli & Ruprecht 2002; He et al. 2002; Gorny et al. 2002; Sharon et al. 2002; Srivastava et al. 2002; Verrier et al. 2001; York et al. 2001; Park et al. 2000; Nyambi et al. 2000; Ly & Stamatatos 2000; Hioe et al.

2000; Grovit-Ferbas et al. 2000; Gorny et al. 2000; Beddows et al. 1999; Hioe et al. 1999; Nyambi et al. 1998; Gorny et al. 1998; Con-

nor et al. 1998; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a; Parren et al. 1998a; Smith et al. 1998; Mondor et al. 1998; Inouye et al. 1998; Ugolini et al. 1997; Gorny

et al. 1997; Hill et al. 1997; Parren et al. 1997b; Boots et al. 1997; Hioe et al. 1997b; Hioe et al. 1997a; Fouts et al. 1997; Binley

et al. 1997a; D'Souza et al. 1997; Sattentau 1996; Trkola et al. 1996a; Jagodzinski et al. 1996; Forthal et al. 1995; Moore & Ho 1995; Moore et al. 1995a; Zolla-Pazner & Sharpe

1995; Zolla-Pazner et al. 1995; Sattentau et al. 1995; Saarloos et al. 1995; Fontenot et al. 1995; Sattentau 1995; Moore et al. 1994a; Gorny et al. 1994; VanCott et al. 1994; Laal et al. 1994; Conley et al. 1994a; Spear et al.

1993; Cavacini et al. 1993a; Keller et al. 1993; Gorny et al. 1993; Karwowska et al. 1992b; Buchbinder et al. 1992; Gorny et al. 1992

Keywords acute/early infection, ADCC, antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain, assay development, assay standardgp160 Antibodies HIV Antibodies Tables

ization/improvement, binding affinity, correceptor, complement, enhancing activity, kinetics, mimotopes, reversion, viral fitness, review, structure, subtype comparisons, Th2, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-neutralization

- 447-52d: 2909 is a human anti-Env NAb that was selected by a neutralization assay and binds to the quaternary structure on the intact virion. ELISA-based competition assays and subsequent mutational analysis determined that the CD4BS and V2 and V3 loops contribute to the 2909 epitope: 2909 binding was inhibited by MAbs 447-52d (anti-V3), 830A (anti-V2), and IgG1b12 (anti-CD4BS) and sCD4. 2909 was not inhibited by MAbs 670, 1418, nor 2G12. Gorny et al. [2005]
- 447-52D: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 447-52D has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)
- 447-52D: Sera from subtype A infected individuals from Cameroon have antibodies that react strongly with subtype A and subtype B V3 loops in fusion proteins, and neutralize SF162 pseudotypes, while sera from 47 subtype B infected individuals reacted only with subtype B. Sera from Cameroon did not neutralize primary A or B isolates, due to indirect masking by the V1/V2 domain rather than due to loss of the target epitope. Neutralization by anti-V3 B clade specific MAbs 447-52D and 4117C was fully blocked by a clade V3 loop fusion protein, but not an A clade fusion protein, while Cameroonian sera neutralization was fully blocked by both A and B clade fusion proteins. Krachmarov et al. [2005] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, subtype comparisons)
- 447-52D: The epitope for the MAb D19 is conserved and embedded in V3. D19 is unique in that for R5 viruses, it was cryptic and did not bind without exposure to sCD4, and for X4 and R5X4 isolates it was constitutively exposed. It had an overlapping binding region with MAbs 447-52D, B4e8, and 268-D, but different reactivity patterns and fine specificity. While B4e8 and 447-52D could bind to the R5 virus BaL in the absence of sCD4, treatment with sCD4 did increase the binding of both B4e8 and 447-52D, but did not impact their ability to neutralize BaL. Lusso *et al.* [2005] (antibody binding site definition and exposure)
- 447-52D: The HIV-1 Bori-15 variant was adapted from the Bori isolate for replication in microglial cells. Bori-15 had increased replication in microglial cells and a robust syncytium-forming phenotype, ability to use low levels of CD4 for infection, and increased sensitivity to neutralization by sCD4 and 17b. Four amino acid changes in gp120 V1-V2 were responsible for this change. Protein functionality and integrity of soluble, monomeric gp120-molecules derived from parental HIV-1 Bori and microglial-adapted HIV-1 Bori-15 was assessed in ELISA binding assays using CD4BS MAbs F105 and IgG1b12, glycan-specific 2G12, and V3-specific 447-52D, and were unchanged.

Association rates of sCD4 and 17b were not changed, but dissociation rates were 3-fold slower for sCD4 and 14-fold slower for 17b. Martín-García *et al.* [2005] (antibody binding site definition and exposure)

- 447-52D: This study is about the V2 MAb C108g, which is type-specific and neutralizes BaL and HXB2. JR-FL is a neutralization resistant strain; modification of JRFL at V2 positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The modification in V2 also increased neutralization sensitivity to V3 MAbs 4117c, 2219, 2191, and 447-52D (447-52D was the only one of the 4 V3 MAbs that could neutralize the unmodified JRFL); but only had minor effects on neutralization by CD4BS MAb 5145A, and broadly neutralizing MAbs IgG1b12, 2G12, and 2F5. Pinter et al. [2005] (antibody binding site definition and exposure)
- 447-52-D: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. V3 MAbs (447-52D, 19b, F245-B4e8 and 39F) bound to the GDMR antigen, but either did not bind or had diminished binding to mCHO. Selvarajah *et al.* [2005] (vaccine-specific epitope characteristics, Th2)
- 447-52D: 93 viruses from different clades were tested for their neutralization cross-reactivity using a panel of HIV antibodies. Neutralization outside of the B clade was very rare, and seemed to depend on the presence of a GPGR V3 tip, which is rare outside of the B clade. Binley et al. [2004] (variant crossrecognition or cross-neutralization, subtype comparisons)
- 447-52D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. The set that can cross-neutralize primary isolates (2182, 2191, 2219, 2412, 2442, 2456) bind V3 but are conformationally sensitive, suggesting some structural conservation despite sequence variation. These MAbs have distinct epitopes relative to 447-52D, a MAb directed at the tip of the V3 loop that also can neutralize many primary isolates. Although 447-52D was selected using a peptide, it has conformational characteristics. Inter-clade cross-neutralization by anti-V3 conformation-dependent MAbs is reduced. Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, review)
- 447-52D: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides, but was an exception in that it is cross-neutralizing. 447-52D neutralized 12/13 clade B viruses. Gorny et al. [2004] (antibody binding site definition and exposure)
- 447-52D: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested

LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of the V3 MAbs 694-98D and 447-52D, that both bind near the tip of the loop, was decreased by both thrombin and trypsin. Ling et al. [2004] (antibody binding site definition and exposure)

- 447-52D: Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The loss of the any of three glycans within or adjacent to the V3 loop (GM299 V3), C2 (GM292 C2), C3 (GM329 C3) increased neutralization susceptibility to 447-52D, but C4 (GM438 C4) or V5 (GM454 V5) removal did not make SF162 more sensitive. V3 glycans tended to shield V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans shielded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004] (antibody binding site definition and exposure, vaccine antigen design)
- 447-52D: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including 447-52D. Pantophlet et al. [2004] (vaccine antigen design)
- 447-52D: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. 5/6 anti-V3 MAbs, including 447-52D, had similar binding affinity to soluble SF162 and JR-FL rgp120s, although the V3 loop differs at three positions (HigpgrafyTtgE for JR-FL and TigpgrafyAtgD for SF162). Pinter et al. [2004] (variant cross-recognition or cross-neutralization)
- 447-52D: A primary isolate, CC1/85, was passaged 19 times in PBMC and gradually acquired increased sensitivity to FAb b12 and sCD4 that was attributed to changes in the the V1V2 loop region, in particular the loss of a potential glycosylation site. The affinity for sCD4 was unchanged in the monomer, suggesting that the structural impact of the change was manifested at the level of the trimer. The passaged virus, CCcon19, retained an R5 phenotype and its neutralization susceptibility to other Abs was essentially the same as CC1/85. 447-52D did not neutralize the primary or passaged variant. Pugach et al. [2004] (reversion, viral fitness, variant cross-recognition or cross-neutralization)

- and diminished cell fusion of H9 cells infected with HIV-1 447-52D: Analysis of the conformation of 447-52D in complex with the V3MN18 peptide (gp12 aa 310-329, KRKRIHIGP-GRAFYTTKN) was undertaken using solid state NMR. The bound peptide had a well defined constrained structure that was in good agreement with solution NMR and crystallographic studies. Sharpe et al. [2004] (structure)
  - 447-52D: The SOS mutant envelope protein introduces a covalent disulfide bond between gp120 surface and gp41 transmembrane proteins into the R5 isolate JR-FL by adding cysteines at residues 501 and 605. Pseudovirions bearing this protein bind to CD4 and co-receptor bearing cells, but do not fuse until treatment with a reducing agent, and are arrested prior to fusion after CD4 and co-receptor engagement. 447-52D was able to neutralize the SOS protein better than the wildtype, but did not neutralize SOS well when added post-attachment, as the V3 loop is involved in co-receptor engagement. Binley et al. [2003] (vaccine antigen design)
  - 447-52D: The Fv fragment (composed of just the light and heavy variable regions, and the smallest intact binding unit of an Ab) of 447-52 D was expressed and purified. Preliminary NMR with the peptide epitope indicates that an NMR structure determination is feasible. Kessler et al. [2003] (antibody sequence, variable domain, structure)
  - 447-52D: This paper attempts to engineer a gp120 molecule that would focus the immune response onto the IgG1b12 epitope. Adding a glycosylation sequon (P313N) to the V3 loop knocked out binding to anti-V3 MAbs loop 2, 19b and 447-52-D. Pantophlet et al. [2003b] (vaccine antigen design)
  - 447-52D: Virion capture assays are not a good predictor of neutralization, and the presentation of epitopes using this assay seems to be different from that of functional Envelope spikes on primary isolates - F105 and b6 could efficiently block the b12-mediated capture of infectious virions in a virus capture, but did not inhibit b12 neutralization - Ab 447-52D was able to potently neutralize 89.6 and to neutralize JR-CSF at a high concentration but poorly neutralized ADA - b12 was potent at neutralizing the three primary virions JR-CSF, ADA, and 89.6, but anti-V3 Abs 447-52D and 19b, which did not neutralize JR-CSF and ADA, captured amounts of p24 equal to or higher than the amounts captured by the neutralizing Ab b12. Poignard et al. [2003] (antibody binding site definition and exposure, assay development, variant cross-recognition or cross-neutralization)
  - 447-52D: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the V3 MAbs used. Zwick et al. [2003] (antibody interactions)
  - 447-52D: Review of NAbs. Ferrantelli & Ruprecht [2002]
  - 447-52D: Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated the six new MAbs all bind to the tip of the V3 loop and cross-

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compete with the MAb 447-52D and are conformationally sensitive – MAbs showed cross-clade binding to native, intact virions of clades A(N=2), B(N=4), and F(N=2), limited binding to C(N=3) and D(N=3), and did not bind to CRF01(subtype E, N=2) – the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay – five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization studies), 654 (anti-CD4BS used as a conformation-sensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control), MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades) – 447-52D bound to primary isolates from all clades except CRF01 (E), was conformationally sensitive and showed the some of the most potent neutralizing activity. Gorny et al. [2002] (variant cross-recognition or cross-neutralization)

- 447-52D: Transgenic mice carrying human genes allowing production of fully human MAbs were used to rapidly create a panel of anti-HIV gp120 MAb producing hybridomas by immunization with HIV SF162 gp120 the previously described human MAbs 5145A(CD4BS), 4117C (plus others, V3) and 697D (and SC258, V2) were used as controls. He *et al.* [2002]
- 447-52D: The feasibility of determining the NMR structure of the V3(MN) peptide bound to the 447-52D Fab fragment was tested and a general strategy for obtaining NMR structures of V3 peptide-Fab fragments developed preliminary NMR spectra for 447-52D complexed to a 23 amino acid V3 peptide was obtained. Sharon *et al.* [2002] (**structure**)
- 447-52D: Oligomeric gp140 (o-gp140) derived from R5 primary isolate US4 was characterized for use as a vaccine reagent—antigen capture ELISA was used to compare the antigenicity of gp120 and o-gp140 using a panel of well characterized MAbs—447-D recognized the gp120 monomer much more readily than o-gp140, suggesting the V3 loop is less exposed on o-gp140 and on intact virions. Srivastava et al. [2002] (antibody binding site definition and exposure, vaccine antigen design)
- 447-52D: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001] (antibody interactions, variant cross-recognition or cross-neutralization)
- 447-52D: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding the dissociation constant, Kd of 447-52D for the cell associated primary and TCLA Envs was equal, 3nM. York et al. [2001] (antibody binding site definition and exposure,

variant cross-recognition or cross-neutralization, binding affinity)

- 447-52D: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer V3 MAbs 447-52D, 838-D, and 1334 bound with a 7-10 fold preference for the oligomer. Gorny *et al.* [2000] (antibody binding site definition and exposure)
- 447-52D: To determine the antigenicity of virus killed by thermal and chemical inactivation, retention of conformation-dependent neutralization epitopes was examined, and exposure of CD4BS epitopes was found to be enhanced (MAbs IgG1b12, 205-46-9, and 205-43-1) binding to 2G12 and 447-52D epitopes was essentially unaltered the 17b CD4i epitope was also exposed. Grovit-Ferbas *et al.* [2000] (vaccine antigen design)
- 447-52D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses – CD4BS MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells – V3 MAbs 447-52-D and 268-10-D did not effect proliferation. Hioe et al. [2000]
- 447-52D: Called 447D SF162 is a neutralization-resistant HIV-1 isolate N-linked glycosylation modifications in the V2 loop of the SF162 gp120 revealed that these sites prevent neutralization by CD4BS MAbs (IgG1b12 and IgGCD4), and protect against neutralization by V3 MAbs (447D and 391-95D) V2-region glycosylation site mutations did not alter neutralization resistance to V2 MAbs (G3.4 and G3.136) or CD4i MAbs (17b and 48d) V2 glycosylation site modification allows infection of macrophages, probably due to glycosylated forms requiring fewer CCR5 molecules for viral entry. Ly & Stamatatos [2000] (antibody binding site definition and exposure)
- 447-52D: A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding V3 MAbs tended to have the most cross-reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H 447-52D showed the highest cross-reactivity, bound to 24/26 viruses tested, but achieved 90% neutralization only against MN, 50% against CA5, and no neutralization was observed for 3 other isolates tested. Nyambi *et al.* [2000] (**subtype comparisons**)
- 447-52D: Six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park *et al.* [2000] (antibody binding site definition and exposure)
- 447-52D: rgp120 derived from a R5X4 subtype B virus, HIV-1 W61D, was used to vaccinate healthy volunteers and the resulting sera were compared with sera from HIV-1 positive subjects and neutralizing MAbs TCLA strains showed enhanced 447-52D neutralization sensitivity relative to PBMC-adapted lines (32X increase between HIV-1 (M2424/PBMC(p0)) and HIV-1 (M2424/H9(p9)) and a >128X increase between HIV-1

dows et al. [1999] (variant cross-recognition or crossneutralization)

- 447-52D: The presence of leukocyte function-associated molecule 1 (LFA-1) promotes virus infectivity and hinders neutralization, and anti-LFA-1 MAbs can enhance the neutralizing effect of anti-HIV V3 MAb 447-52D and anti-HIV CD4BS MAb IgG1b12 - non-neutralizing anti-HIV CD4BS MAb 654-D did not become neutralizing in the presence of anti-LFA-1 MAbs. Hioe *et al.* [1999]
- 447-52D: Review of clade specificity and anti-V3 HIV-1-Abs. Zolla-Pazner *et al.* [1999a] (**review**, **subtype comparisons**)
- 447-52D: MAb peptide-reactivity pattern clustered with the immunological related MAbs: 1334, 419, 504, 447, 453 and 537 - the core amino acids GP tended to be critical for reactivity in this group – 447 reacted with peptides containing GPGR, but also with many lacking this sequence (GPGQ, for example), and it failed to react with 2/14 peptides containing GPGR, illustrating the importance of context. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 447-52D: Ab from gp120 vaccinated individuals prior to infection, who subsequently became HIV infected, could not achieve 90% neutralization of the primary virus by which the individuals were ultimately infected - these viruses were not particularly refractive to neutralization, as determined by their susceptibility to neutralization by MAbs 2G12, IgG1b12, 2F5 and 447-52D. Connor et al. [1998]
- 447-52D: Kinetic parameters were measured, and the association rates were similar, but dissociation rate constants were quite variable for V3 MAbs, 1324E was comparable to 447-52D. Gorny *et al.* [1998] (**kinetics**)
- 447-52D: Called 447-D 447-D resistance took longer to acquire in virus with the M184V substituted RT, and had the form (AAC N to TAC Y) at position 5 of the V3 loop, rather than the GPGR to GPGR resistance found with wildtype RT. Inouye et al. [1998]
- 447-52D: Inhibits binding of Hx10 to both CD4 positive and negative HeLa cells. Mondor et al. [1998] (variant crossrecognition or cross-neutralization)
- 447-52D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H - 447-52D was the most potent and cross-reactive of 18 human MAbs tested and was the only MAb which bound to virions from isolates CA20 (subtype F), CA13 (subtype H), and VI526 (subtype G) Nyambi et al. [1998] (subtype comparisons)
- 447-52D: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated - authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a] (antibody binding site definition and exposure)
- 447-52D: Called 447-52-D The tip of the MN V3 loop was inserted into cold causing human rhinovirus 14 (HRV14) chimeras were immunoselected, and chimeric viruses were neutralized by anti-V3 loop antibodies, and 447-52D was among the Abs used - chimeric viruses elicited potent NAbs in guinea pigs against ALA-1 and MN. Smith et al. [1998] (vaccine antigen design)

- (W61D/PBMC) and HIV-1 (W61D/SupT1) isolates) Bed- 447-52D: Abs that recognize discontinuous epitopes can identify mimotopes from a phage peptide display library – 447-52D has an epitope involving the tip of the V3 loop, that was previously studied with this method Keller et al. [1993] – in Keller et al., with no competition, LxGPxR was the most common six-mer, 38% of the peptides – after competition with a gp120 IIIB ligand (QRGPGR)i, RGPxR was the most common and one peptide had the sequence ORGPGR, showing type specific mimotopes can be enriched by strain specific ligand competition protocols Boots et al. [1997]. Boots et al. [1997]; Keller et al. [1993] (antibody binding site definition and exposure, mimotopes)
  - 447-52D: In a multilaboratory blinded study, failed to consistently neutralize any of nine B clade primary isolates – many of these isolates had the GPGR motif at the apex of the V3 loop. D'Souza et al. [1997] (variant cross-recognition or cross-neutralization, assay standardization/improvement)
  - 447-52D: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding – 447-52D bound monomer, oligomer, and neutralized JRFL. Fouts et al. [1997] (antibody binding site definition and exposure)
  - 447-52D: Used as a control for comparison to five V3 RF selected antibodies - 447-52D was reactive with A, B, and C clade peptides, but not E. Gorny et al. [1997] (subtype comparisons)
  - 447-52D: Called 447 gp120 can inhibit MIP-1alpha from binding to CCR5, but this inhibitory effect is blocked by preincubation of gp120 with three anti-V3 MAbs: 447, 257, 1027 - MAb 670 which binds in the C5 region had no effect. Hill et al. [1997] (co-receptor)
  - 447-52D: Tested using a resting cell neutralization assay. Hioe et al. [1997a] (assay standardization/improvement)
  - 447-52D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs – BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) - isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations – US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe et al. [1997b] (variant cross-recognition or cross-neutralization)
  - 447-52D: Neutralizes TCLA strains but not primary isolates. Parren et al. [1997b] (variant cross-recognition or crossneutralization)
  - 447-52D: Viral binding inhibition by 447-D was correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5) Ugolini et al. [1997] (antibody binding site definition and exposure)
  - 447-52D: Called 447-52-D The sulfated polysaccharide curdlan sulfate (CRDS) binds to the Envelope of T-tropic viruses and neutralizes virus - CRDS inhibits 447-52D binding. Jagodzinski et al. [1996] (antibody binding site definition and exposure)

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- 447-52D: Review: called 447-52-D only four epitopes have been described which can stimulate a useful neutralizing response to a broad spectrum of primary isolates, represented by the binding sites of MAbs: 447-52-D, 2G12, Fab b12, and 2F5. Sattentau [1996] (variant cross-recognition or crossneutralization, review)
- 447-52D: Neutralizes JR-FL strongly inhibits gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a] (co-receptor, variant cross-recognition or cross-neutralization)
- 447-52D: Called 447 The tip of the V3 loop was presented in a mucin backbone – higher valency correlates with stronger affinity constant. Fontenot *et al.* [1995] (vaccine antigen design)
- 447-52D: Neutralizing (- complement), no ADCC activity, and no viral enhancing activity. Forthal *et al.* [1995] (ADCC, complement, enhancing activity)
- 447-52D: Binding affected by identity of amino acids flanking GPGR core – poor breadth of primary virus neutralization. Moore *et al.* [1995a] (variant cross-recognition or cross-neutralization)
- 447-52D: Review: the V3 loop motif GPGR is not common outside subtype B isolates, MAb 19b is more cross-reactive than 447-52D. Moore & Ho [1995] (variant cross-recognition or cross-neutralization)
- 447-52D: Ab-mediated activation of complement on HIV+cells is higher than Ab independent activation—what has been termed "Ab independent" in fact results in part from IgM in normal human serum that is HIV-cross-reactive. Saarloos *et al.* [1995] (complement)
- 447-52D: Called 447d Formalin inactivation of virus at 0.1% formalin for 10 hours at 4 degrees was optimal for inactivation of virus while maintaining epitope integrity. Sattentau *et al.* [1995] (vaccine antigen design)
- 447-52D: Serotyping study using flow-cytometry bound only to GPGR V3 loop tips. Zolla-Pazner *et al.* [1995] (antibody binding site definition and exposure)
- 447-52D: Neutralization of primary and prototype laboratory HIV-1 isolates using a resting cell assay enhances sensitivity.
   Zolla-Pazner & Sharpe [1995] (assay development, variant cross-recognition or cross-neutralization)
- 447-52D: Requires GPxR at the tip of the V3 loop, common in B clade neutralized primary isolates. Conley *et al.* [1994a] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 447-52D: Mild oxidation of carbohydrate moieties does not alter binding. Gorny et al. [1994] (antibody binding site definition and exposure)
- 447-52D: Neutralization synergy in combination with CD4 binding domain MAbs. Laal *et al.* [1994] (antibody interactions)
- 447-52D: Competition studies with human sera from seroconverting individuals showed that anti-CD4 BS antibodies can arise very early in infection, comparable or prior to anti-V3 antibodies. Moore *et al.* [1994a] (acute/early infection)
- 447-52D: GPGQ in MAL resulted in enhanced dissociation GPGQ in CM234 or K14T did not bind binding affected by identity of amino acids flanking GPGR core. VanCott *et al.* [1994] (antibody binding site definition and exposure)

- 447-52D: Additive neutralization of MN and SF2 when combined with CD4 binding site MAb F105 supra-additive neutralization of RF. Cavacini *et al.* [1993a] (antibody interactions)
- 447-52D: Neutralizes MN and IIIB: GPGR, and binds SF2: GPGR. Gorny et al. [1993] (variant cross-recognition or cross-neutralization)
- 447-52D: Peptide phage library showed that any of the residues ADGLMNQRS in the X position tolerated in peptides that react well with the antibody. Keller et al. [1993] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 447-52D: Complement mediated virolysis of IIIB, but not in the presence of sCD4. Spear *et al.* [1993] (**complement**)
- 447-52D: 60-fold increase in neutralization potency when combined 1:1 with human MAb 588-D. Buchbinder *et al.* [1992] (antibody interactions)
- 447-52D: Requires GPXR at the tip of the V3 loop neutralizes a broad array of B clade lab isolates. Gorny *et al.* [1992] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 447-52D: Reacts with MN, NY5, CDC4, SF2, RF, WM52, and HXB2. Karwowska *et al.* [1992b] (variant cross-recognition or cross-neutralization)

No. 745

MAb ID C8

HXB2 Location gp160 (727–732)

Author Location gp41 (727–732 BH10)

**Epitope** PDRPEG

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG1)

**Ab Type** C-term

**References** Heap *et al.* 2005a; Dimmock 2005; McLain *et al.* 2001; Abacioglu *et al.* 1994; Pincus *et al.* 

1993; Pincus & McClure 1993

Keywords review

- C8: This review summarizes the complex antigenic properties
  of an external loop in the gp41 tail (spanning the Kennedy
  sequence), highlighting specific MAbs. C8 binds to the epitope
  PDRPEG and does not neutralize virus. Dimmock [2005]
  (review)
- C8: Unlike SAR1, a MAb that binds near the C8 epitope within the Kennedy peptide, C8 cannot inhibit fusion between HIV-1 infected and target cells. C8 recognizes PDRPEG on the surface of HIV-1 infected cells, but not on virions and is non-neutralizing. Heap et al. [2005a]
- C8: The substitution 725 RG (P[R->G]GPDRPEGIEEEGGERDRDRS) alters the antigenic exposure of this region on the virion resulting in the loss of the downstream neutralizing epitope ERDRD, increased exposure of the epitope GPDRPEG in the virion, while the epitope IEEE remains unchanged. McLain *et al.* [2001]
- C8: Epitope boundaries mapped by peptide scanning. Abacioglu et al. [1994]

• C8: Immunotoxin of C8 coupled to ricin-A does not mediate cells killing, and is not affected by sCD4. Pincus & McClure

• C8: Ab response in IIIB lab workers was compared to gp160 LAI vaccine recipients - C8 was used as a control - the dominant response among vaccinees was to this mid-gp41 region, but not among the infected lab workers - Abs binding this region do not neutralize, bind to infected cells, nor serve as immunotoxins. Pincus et al. [1993]

**No.** 746

MAb ID B31

HXB2 Location gp160 (727-734)

Author Location gp41 (727–734 BH10)

Epitope PDRPEGIE

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI

HIV component: gp160

Species (Isotype) mouse (IgG1)

References Abacioglu et al. 1994

• B31: Epitope boundaries mapped by peptide scanning. Abacioglu et al. [1994]

No. 747

MAb ID B33

HXB2 Location gp160 (727-734)

Author Location gp41 (727–734 BH10)

Epitope PDRPEGIE

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43

HIV component: gp160

Species (Isotype) mouse (IgG1)

References Bristow et al. 1994; Abacioglu et al. 1994

- B33: Epitope boundaries mapped by peptide scanning IgG1. Species (Isotype) mouse Abacioglu et al. [1994]
- B33: There are two MAbs in the literature named B33, see also gp120, positions 123-142 - MAb generated in a study of the humoral immune response to Baculovirus-expressed mis-folded rgp160 IIIB:NL43, MicroGenSys. Bristow et al. [1994]

**No.** 748

**MAb ID** 1576

**HXB2 Location** gp160 (728–745)

Author Location gp41 (735–752 IIIB)

Epitope DRPEGIEEEGGERDRDRS

Neutralizing no

Immunogen vaccine

Vector/Type: poliovirus Strain: B clade IIIB

HIV component: gp41

Species (Isotype) mouse

References Vella et al. 1993

• 1576: Not neutralizing. Vella et al. [1993]

**No.** 749

**MAb ID** 1578

**HXB2 Location** gp160 (728–745)

Author Location gp41 (735–752 IIIB)

Epitope DRPEGIEEEGGERDRDRS

Neutralizing no

Immunogen vaccine

Vector/Type: poliovirus Strain: B clade IIIB

HIV component: gp41

**Species (Isotype)** mouse

References Vella et al. 1993; Evans et al. 1989

- 1578: Core epitope: IEEE in this study, neutralized IIIB, but not RF or MN. Vella et al. [1993]
- 1578: No neutralizing activity epitope may be formed by regions from both poliovirus and HIV. Evans et al. [1989]

**No.** 750

**MAb ID** 1579

HXB2 Location gp160 (728–745)

Author Location gp41 (735–752 IIIB)

Epitope DRPEGIEEEGGERDRDRS

Neutralizing no

Immunogen vaccine

Vector/Type: poliovirus Strain: B clade IIIB

HIV component: gp41

Species (Isotype) mouse

References Vella et al. 1993

• 1579: Core epitope: IEEE - neutralized IIIB, but not RF or MN. Vella et al. [1993]

No. 751

**MAb ID** 1583

**HXB2 Location** gp160 (728–745)

Author Location gp41 (735–752 IIIB)

Epitope DRPEGIEEEGGERDRDRS

Neutralizing no

Immunogen vaccine

Vector/Type: poliovirus Strain: B clade IIIB

HIV component: gp41

References Heap et al. 2005a; Dimmock 2005; Sattentau

et al. 1995; Vella et al. 1993; Evans et al. 1989

Keywords review

- 1583: This review summarizes the complex antigenic properties of an external loop in the gp41 tail (Kennedy sequence), highlighting specific MAbs. 1577 and 1583 bind to the epitope ERDRD and do not neutralize virus. Dimmock [2005]
- 1583: Unlike SAR1, a MAb that binds near the 1583 epitope within the Kennedy peptide, 1583 cannot inhibit fusion between HIV-1 infected and target cells. 1583 and 1577 neturalize only in the presence of complement. Heap et al. [2005a]
- 1583: Suggested to bind to a cytoplasmic domain, epitope not exposed at the surface of HIV-1 infected cells. Sattentau et al.
- 1583: Core epitope: ERDRD Could neutralize HIV IIIB but not HIV RF. Vella et al. [1993]
- 1583: Neutralizing activity, less broad than 1577. Evans et al. [1989]

**No.** 752

**MAb ID** 1899

HXB2 Location gp160 (728–745) Author Location gp41 (735-752 IIIB) gp160 Antibodies **HIV Antibodies Tables** 

Epitope DRPEGIEEEGGERDRDRS Vector/Type: peptide Strain: B clade IIIB Neutralizing no HIV component: gp41 Immunogen vaccine **Species (Isotype)** mouse ( $IgM\kappa$ ) Vector/Type: poliovirus Strain: B clade IIIB References Dalgleish et al. 1988; Mani et al. 1994 • 41-1: This antibody gp41(735-752 IIIB) Dalgleish et al. [1988] HIV component: gp41 Species (Isotype) mouse seems to have been named the same as a different MAb to References Vella et al. 1993 gp41(584-609) Mani et al. [1994]. Dalgleish et al. [1988]; • 1899: Could neutralize HIV IIIB and HIV RF. Vella et al. Mani et al. [1994] • 41-1: Neutralizes HIV-1 but not HIV-2 strains. Dalgleish et al. [1993] **No.** 753 **MAb ID** 1907 No. 757 HXB2 Location gp160 (728-745) **MAb ID** 41-2 Author Location gp41 (735-752 IIIB) HXB2 Location gp160 (728–745) **Epitope** DRPEGIEEEGGERDRDRS Author Location gp41 (735–752 IIIB) Neutralizing no Epitope DRPEGIEEEGGERDRDRS Neutralizing no Immunogen vaccine Immunogen vaccine Vector/Type: poliovirus Strain: B clade IIIB HIV component: gp41 Vector/Type: peptide Strain: B clade IIIB Species (Isotype) mouse HIV component: gp41 References Vella et al. 1993 **Species (Isotype)** mouse ( $IgM\kappa$ ) • 1907: Could not neutralize HIV IIIB, RF or MN. Vella et al. References Dalgleish et al. 1988 [1993] • 41-2: Neutralizes HIV-1 but not HIV-2 strains. Dalgleish et al. [1988] **No.** 754 **MAb ID** 1908 No. 758 HXB2 Location gp160 (728-745) **MAb ID** 41-3 Author Location gp41 (735–752 IIIB) HXB2 Location gp160 (728–745) Author Location gp41 (735–752 IIIB) Epitope DRPEGIEEEGGERDRDRS Epitope DRPEGIEEEGGERDRDRS Neutralizing no Immunogen vaccine Neutralizing no Vector/Type: poliovirus Strain: B clade IIIB Immunogen vaccine HIV component: gp41 Vector/Type: peptide Strain: B clade IIIB HIV component: gp41 Species (Isotype) mouse References Sattentau et al. 1995; Vella et al. 1993; Evans **Species (Isotype)** mouse ( $IgM\kappa$ ) et al. 1989 References Dalgleish et al. 1988 • 1908: Cytoplasmic domain, epitope not exposed at the surface • 41-3: Neutralizes HIV-1 but not HIV-2 strains. Dalgleish et al. of HIV-1 infected cells. Sattentau et al. [1995] [1988] • 1908: Neutralized IIIB, but not RF or MN. Vella et al. [1993] No. 759 No. 755 MAb ID ED6 **MAb ID** 1909 HXB2 Location gp160 (728-745) HXB2 Location gp160 (728-745) Author Location gp41 (735–752 IIIB) Author Location gp41 (735-752 IIIB) Epitope DRPEGIEEEGGERDRDRS Epitope DRPEGIEEEGGERDRDRS Neutralizing no Neutralizing no Immunogen Immunogen vaccine Species (Isotype) mouse (IgM) Vector/Type: poliovirus Strain: B clade IIIB References Evans et al. 1989 HIV component: gp41 No. 760 Species (Isotype) mouse **MAb ID** LA9 (121-134) References Vella et al. 1993 **HXB2 Location** gp160 (728–745) • 1909: Neutralized HIV IIIB but not HIV RF. Vella et al. [1993] Author Location gp41 (735–752 IIIB) **No.** 756 **Epitope** DRPEGIEEEGGERDRDRS **MAb ID** 41-1 Neutralizing no HXB2 Location gp160 (728–745) Immunogen Author Location gp41 (735–752 IIIB) Species (Isotype) mouse (IgM) Epitope DRPEGIEEEGGERDRDRS References Evans et al. 1989 Neutralizing no No. 761

Immunogen vaccine

**MAb ID** 1575

HXB2 Location gp160 (728-745)

Author Location gp41 (735-752 IIIB)

Epitope DRPEGIEEEGGERDRDRS

Neutralizing no

Immunogen vaccine

Vector/Type: poliovirus Strain: B clade IIIB

HIV component: gp41

Species (Isotype) mouse

Ab Type C-term

Research Contact C. Vella, NIBSC, Potters Bar UK

**References** Heap *et al.* 2005a; Dimmock 2005; Cleveland *et al.* 2000a; Buratti *et al.* 1997; Vella *et al.* 

1993; Evans et al. 1989

Keywords review

- 1575: This review summarizes the complex antigenic properties of an external loop in the gp41 tail (Kennedy sequence), highlighting specific MAbs. 1575 is noted to bind to the epitope IEEE and does not neutralize virus. Dimmock [2005] (review)
- 1575: Unlike SAR1, a MAb that binds near the 1575 epitope within the Kennedy peptide, 1575 cannot inhibit fusion between HIV-1 infected and target cells. Heap *et al.* [2005a]
- 1575: Ab binding to IEEE suppresses neutralizing Ab binding to adjacent epitope ERDRD. Cleveland *et al.* [2000a]
- 1575: Study shows that MAb 1575 can recognize the IEEE sequence in both gp41, and in the HPG30 region of the p17 protein motif is conserved in both regions in different HIV-1 clades. Buratti *et al.* [1997]
- 1575: Core epitope: IEEE neutralized IIIB, but not RF or MN. Vella et al. [1993]
- 1575: Neutralizing activity, less broad than 1577. Evans *et al.* [1989]

No. 762

**MAb ID** 88-158/02

HXB2 Location gp160 (732–747)

Author Location gp41 (732–752 IIIB)

Epitope GIEEEGGERDRDRSIR

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp41

Species (Isotype) mouse (IgG2b)

References Niedrig et al. 1992a

• 88-158/02: Mild inhibition of *in vitro* activity at high MAb concentrations – profound enhancing activity at low concentrations – significant reactivity to virion – domain non-immunogenic in humans. Niedrig *et al.* [1992a]

**No.** 763

MAb ID 88-158/022

**HXB2 Location** gp160 (732–747)

Author Location gp41 (732–752 IIIB)

Epitope GIEEEGGERDRDRSIR

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp41

Species (Isotype) mouse (IgG2b)

References Niedrig et al. 1992a

88-158/022: Mild inhibition of *in vitro* activity at high MAb concentrations – profound enhancing activity at low concentrations – significant reactivity to virion – domain non-immunogenic in humans. Niedrig *et al.* [1992a]

No. 764

**MAb ID** 88-158/079

**HXB2 Location** gp160 (732–747)

Author Location gp41 (732-752 IIIB)

Epitope GIEEEGGERDRDRSIR

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp41

Species (Isotype) mouse (IgG1)

References Niedrig et al. 1992a

88-158/079: Mild inhibition of HIV in vitro at high MAb concentrations – profound enhancing activity at low concentrations

 weak binding to virion – domain non-immunogenic in humans.
 Niedrig et al. [1992a]

**No.** 765

MAb ID polyclonal

**HXB2 Location** gp160 (733–736)

Author Location gp41 (735–752 IIIB)

Epitope IEEE

**Neutralizing** L

Immunogen vaccine

Vector/Type: Cowpea mosaic virus HIV

component: gp41

Species (Isotype) mouse (IgG)

Ab Type C-term

References McLain et al. 2001; Cleveland et al. 2000b

- The substitution 725 RG (P[R—>G]GPDRPEGIEEEGGERDRDRS) alters the antigenic exposure of this region on the virion resulting in the loss of the downstream neutralizing epitope ERDRD, increased exposure of the epitope GPDRPEG in the virion, while the epitope IEEE remains unchanged. McLain *et al.* [2001]
- When PRGPDRPEGIEEEGGERDRDRS was used as antigen an immunodominant, non-neutralizing response to IEEE was observed, but immunization GERDRDR shifts the response to ERDRD. Cleveland et al. [2000b]

**No.** 766

MAb ID polyclonal

HXB2 Location gp160 (733–736)

Author Location gp41 (735–752 NL43)

Epitope IEEE

**Neutralizing** L

Immunogen vaccine

Vector/Type: Cowpea mosaic virus HIV

component: gp41

 $\textbf{Species (Isotype)} \ \ \text{mouse (IgG)}$ 

Ab Type C-term

References McLain et al. 2001

• The 725 RG (P[Rsubstitution >GIGPDRPEGIEEEGGERDRDRS) alters the antigenic exposure of this region on the virion resulting in the loss of the downstream neutralizing epitope ERDRD, increased exposure of the epitope GPDRPEG in the virion, while the epitope IEEE remains unchanged. McLain et al. [2001]

> No. 767 MAb ID B8

HXB2 Location gp160 (733-741) Author Location gp41 (733–741 BH10)

Epitope IEEEGGERD

Neutralizing no Immunogen vaccine

> Vector/Type: protein Strain: B clade LAI HIV component: gp160

Species (Isotype) mouse (IgG1)

References Abacioglu et al. 1994; Pincus et al. 1993

- B8: Epitope boundaries mapped by peptide scanning. Abacioglu et al. [1994]
- B8: Ab response in IIIB lab workers was compared to gp160 LAI vaccine recipients - B8 was used as a control - the dominant response among vaccinees was to this mid-gp41 region, but not among the infected lab workers - Abs binding this region do not neutralize, bind to infected cells, nor serve as immunotoxins. Pincus et al. [1993]

**No.** 768

MAb ID SAR1

**HXB2 Location** gp160 (738–744)

Author Location gp41 (735–752 IIIB)

Epitope GERDRDR

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: Cowpea mosaic virus HIV component: gp41

**Species** (**Isotype**) mouse ( $IgG2a\kappa$ )

Ab Type C-term

References Heap et al. 2005a; Dimmock 2005

Keywords antibody binding site definition and exposure, review

- SAR1: This ERDRD-specific MAb recognizes an externalized loop of the gp41 C-terminal domain, and can reduce the yield of infectious progeny and inhibit fusion post-attachment, but not neutralize free virus. Dimmock [2005] (review)
- SAR1: This paper confirms the post-attachment neutralization (PAN) activity of this gp41 C-terminal tail-specific Ab – cell fusion between infected and uninfected cells is inhibited and is temperature dependent. This MAb does not neutralize free virus, and due to PAN activity, is considered to bind an epitope on the external surface of the membrane. The SAR1 epitope is exposed optimally after infected and non-infected cells have attached, prior to fusion. sCD4 binding does not enhance binding of SAR1. MAbs to adjacent epitopes do not have PAN activity. Heap et al. [2005a] (antibody binding site definition and exposure)

**No.** 769 **MAb ID** 1577 HXB2 Location gp160 (739–743)

Author Location gp41 (735–752 IIIB)

**Epitope** ERDRD

Neutralizing no Immunogen vaccine

Vector/Type: poliovirus Strain: B clade IIIB

HIV component: gp41

Species (Isotype) mouse

**Ab Type** C-term

Research Contact C. Vella or Morag Ferguson (NIBSC, Potters

Bar UK)

References Heap et al. 2005a; Dimmock 2005; Cleveland

et al. 2000a; Vella et al. 1993; D'Souza et al.

1991; Evans et al. 1989

Keywords review

- 1577: UK Medical Research Council AIDS reagent: ARP317.
- 1577: NIH AIDS Research and Reference Reagent Program:
- 1577: This review summarizes the complex antigenic properties of an external loop in the gp41 tail (Kennedy sequence), highlighting specific MAbs. 1577 and 1583 bind to the epitope ERDRD and do not neutralize virus. Dimmock [2005] (review)
- 1577: Unlike SAR1, a MAb that binds near the 1577 epitope within the Kennedy peptide, 1577 cannot inhibit fusion between HIV-1 infected and target cells. Heap et al. [2005a]
- 1577: Ab binding to IEEE suppresses neutralizing Ab binding to adjacent epitope ERDRD. Cleveland et al. [2000a]
- 1577: Core epitope: ERDRD could neutralize HIV IIIB and HIV RF. Vella et al. [1993]
- 1577: Non-neutralizing in this multi-lab study. D'Souza et al. [1991]
- 1577: Raised against IIIB peptide chimera neutralized African and American HIV-1 lab strains. Evans et al. [1989]

**No.** 770

MAb ID polyclonal (EPES)

HXB2 Location gp160 (739–743)

Author Location gp41 (735–752 IIIB)

**Epitope** ERDRD

Subtype B

Neutralizing L

Immunogen vaccine

Vector/Type: Cowpea mosaic virus HIV

component: gp41

Species (Isotype) mouse (IgG1, IgG2a, IgG2b)

Ab Type C-term

References Dimmock 2005; McLain et al. 2001; Cleve-

land et al. 2000b

Keywords review

• ERDRD-specific IgG recognizes an externalized loop of the gp41 C-terminal domain, and these polyclonal antibodies are the only ones known to bind to this domain that can neutralize cell-free virus. This paper calls these antibodies EPES for Epitope Purified and ERDRD specific. Dimmock [2005] (review)

• The substitution 725 RG (P[R->GIGPDRPEGIEEEGGERDRDRS) alters the antigenic exposure of this region on the virion resulting in the loss of the downstream neutralizing epitope ERDRD, increased exposure of the epitope GPDRPEG in the virion, while the epitope IEEE remains unchanged. McLain et al. [2001]

• ERDRD-specific IgG recognizes an externalized loop of the gp41 C-terminal tail with high affinity – neutralized HIV-1 B clade strains IIIB, NL-4.3, RF, MN and D clade virus CBL-4, but HXB-2D (clade B) was not recognized - when PRGP-DRPEGIEEEGGERDRDRS was used as antigen an immunodominant, non-neutralizing response to IEEE was observed, but immunization GERDRDR shifts the response to ERDRD -NAb does not inhibit attachment of free virus, but does inhibit Research Contact G. by an event that precedes fusion-entry. Cleveland et al. [2000b]

No. 771

MAb ID DZ

HXB2 Location gp160 (822-855) Author Location gp41 (827–860 BRU)

Neutralizing L Immunogen vaccine

> Vector/Type: vaccinia Strain: B clade IIIB HIV component: Env

**Species (Isotype)** human (IgG1 $\lambda$ ) References Boyer et al. 1991

• DZ: Weakly neutralizing IIIB - binds to peptides 827-843 and 846-860 of BRU - reacted specifically with IIIB and RF. Boyer et al. [1991]

**No.** 772

**MAb ID** 1010

HXB2 Location gp160

**Author Location** gp41

**Epitope** 

Subtype B

Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human

**Ab Type** gp41 six-helix bundle and the internal trimeric **Research Contact** G. coiled-coil of N-helices

Research Contact G.

Marius Clore or Carole Bewley, NIH, Bethesda, Maryland. marius@intra.niddk.nih.gov or caroleb@mail.nih.gov

References Louis et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation

• 1010: This is one of nine anti-gp41 bivalent Fabs selected from a non-immune human phage display library. This Fab was classified as class B, and inhibits LAV mediated Env-mediated cell fusion with an IC50 of 6 +/-2 ug/ml (the range for the 9 new Fabs was 6-61 ug/ml - for context, 2F5 and 2G12 (IC50s of 0.5-1.5 ug/ml) were about an order of magnitude more potent in this assay than the best Fabs generated here). 1010 and 1018, both class B, were the most potent Fabs. The class B Fabs interact with the six-helix bundle and the internal coiled-coil

of N-helices of gp41. Louis et al. [2005] (antibody binding site definition and exposure, antibody generation)

No. 773

**MAb ID** 1014

HXB2 Location gp160

**Author Location** gp41

**Epitope** 

Subtype B

Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human

Ab Type gp41six-helix bundle

Marius Clore

Carole Be-NIH. Bethesda. wlev. Maryland. marius@intra.niddk.nih.gov or car-

or

oleb@mail.nih.gov

References Louis et al. 2005

Keywords antibody binding site definition and exposure, antibody generation

Epitope VAEGTDRVIEVVQGACRAIRHIPRRIRQGLER- • 1014: This is one of nine anti-gp41 bivalent Fabs selected from a non-immune human phage display library. This Fab was classified as class A, and inhibits LAV mediated Env-mediated cell fusion with an IC50 of 36 +/-1 ug/ml (the range for the 9 new Fabs was 6-61 ug/ml - for context, 2F5 and 2G12 were about an order of magnitude more potent in this assay than the best Fabs generated here). Class A Fabs interact with the sixhelix bundle of gp41. Louis et al. [2005] (antibody binding site definition and exposure, antibody generation)

No. 774

**MAb ID** 1018

HXB2 Location gp160

**Author Location** gp41

**Epitope** 

Subtype B

Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human

Ab Type gp41 six-helix bundle and the internal trimeric coiled-coil of N-helices

Marius Clore or Carole Bewley, NIH, Bethesda, Maryland.

marius@intra.niddk.nih.gov car-

oleb@mail.nih.gov

References Louis et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation

1018: This is one of nine anti-gp41 bivalent Fabs selected from a non-immune human phage display library. This Fab was classified as class B, and inhibits LAV mediated Env-mediated cell fusion with an IC50 of 7 +/- 1 ug/ml (the range for the 9 new Fabs was 6-61 ug/ml - for context, 2F5 and 2G12 (IC50s of 0.5-1.5 ug/ml) were about an order of magnitude more potent in this assay than the best Fabs generated here). 1010 and 1018, both class B, were the most potent Fabs. The class B Fabs interact with the six-helix bundle and the internal coiled-coil of N-helices of gp41. Louis et al. [2005] (antibody binding site definition and exposure, antibody generation)

No. 775 **MAb ID** 1019 HXB2 Location gp160 **Author Location** gp41 **Epitope** Subtype B **Neutralizing** Immunogen in vitro stimulation or selection Species (Isotype) human **Ab Type** gp41six-helix bundle

Research Contact G. Clore Be-Marius Carole or NIH, wley, Bethesda. Maryland. marius@intra.niddk.nih.gov oleb@mail.nih.gov

References Louis et al. 2005

Keywords antibody binding site definition and exposure, antibody generation

• 1019: This is one of nine anti-gp41 bivalent Fabs selected from a non-immune human phage display library. This Fab was classified as class A, and inhibits LAV mediated Env-mediated cell fusion with an IC50 of 61 +/- 20 ug/ml (the range for the 9 new Fabs was 6-61 ug/ml - for context, 2F5 and 2G12 were about an order of magnitude more potent in this assay than the best Fabs generated here). Class A Fabs interact with the sixhelix bundle of gp41. Louis et al. [2005] (antibody binding site definition and exposure, antibody generation)

No. 776 **MAb ID** 1020 HXB2 Location gp160 Author Location gp41 **Epitope** Subtype B **Neutralizing** 

Immunogen in vitro stimulation or selection

Species (Isotype) human

**Ab Type** gp41 six-helix bundle and the internal trimeric coiled-coil of N-helices

Research Contact G. Marius Clore Carole Bewley, NIH, Bethesda, Maryland. marius@intra.niddk.nih.gov caroleb@mail.nih.gov

References Louis et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation

• 1020: This is one of nine anti-gp41 bivalent Fabs selected from a non-immune human phage display library. This Fab was classified as class B, and inhibits LAV mediated Env-mediated cell fusion with an IC50 of 20 +/- 3 ug/ml (the range for the 9 new Fabs was 6-61 ug/ml – for context, 2F5 and 2G12 (IC50s of 0.5-1.5 ug/ml) were about an order of magnitude more potent in this assay than the best Fabs generated here). The class B Fabs interact with the six-helix bundle and the internal coiled-coil of N-helices of gp41. Louis et al. [2005] (antibody binding site definition and exposure, antibody generation)

**No.** 777 **MAb ID** 1022 HXB2 Location gp160 **Author Location** gp41

**Epitope** Subtype B **Neutralizing** 

Immunogen in vitro stimulation or selection

Species (Isotype) human

**Ab Type** gp41 six-helix bundle and the internal trimeric coiled-coil of N-helices

Research Contact G. Marius Clore Carole Beor wley, NIH, Bethesda, Maryland. marius@intra.niddk.nih.gov caroleb@mail.nih.gov

References Louis et al. 2005

Keywords antibody binding site definition and exposure, antibody generation

• 1022: This is one of nine anti-gp41 bivalent Fabs selected from a non-immune human phage display library. This Fab was classified as class B, and inhibits LAV mediated Env-mediated cell fusion with an IC50 of 40 +/- 10 ug/ml (the range for the 9 new Fabs was 6-61 ug/ml - for context, 2F5 and 2G12 (IC50s of 0.5-1.5 ug/ml) were about an order of magnitude more potent in this assay than the best Fabs generated here). The class B Fabs interact with the six-helix bundle and the internal coiled-coil of N-helices of gp41. Louis et al. [2005] (antibody binding site definition and exposure, antibody generation)

No. 778 **MAb ID** 1034 HXB2 Location gp160 Author Location gp41 **Epitope** Subtype B Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human

Ab Type gp41 internal trimeric coiled-coil of N-helices Research Contact G. Marius Clore or Carole Bewley, NIH, Bethesda, Maryland. marius@intra.niddk.nih.gov caroleb@mail.nih.gov

References Louis et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation

• 1034: This is one of nine anti-gp41 bivalent Fabs selected from a non-immune human phage display library. This Fab was classified as class C, and inhibits LAV mediated Env-mediated cell fusion with an IC50 of 17 +/- 2 ug/ml (the range for the 9 new Fabs was 6-61 ug/ml - for context, 2F5 and 2G12 (IC50s of 0.5-1.5 ug/ml) were about an order of magnitude more potent in this assay than the best Fabs generated here). The class C Fabs interact with the internal coiled-coil of N-helices of gp41. Louis et al. [2005] (antibody binding site definition and exposure, antibody generation)

No. 779 **MAb ID** 1492 HXB2 Location gp160 **Author Location** gp41 **Epitope** Subtype B Neutralizing

Immunogen in vitro stimulation or selection

Species (Isotype) human

**Ab Type** gp41 internal trimeric coiled-coil of N-helices Clore Research Contact G. Marius or Carole Be-NIH, Bethesda, Maryland. wley, marius@intra.niddk.nih.gov caroleb@mail.nih.gov

References Louis et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation

• 1492: This is one of nine anti-gp41 bivalent Fabs selected from a non-immune human phage display library. This Fab was classified as class C, and inhibits LAV mediated Env-mediated cell fusion with an IC50 of 25 +/- 2 ug/ml (the range for the 9 new Fabs was 6-61 ug/ml - for context, 2F5 and 2G12 (IC50s of 0.5-1.5 ug/ml) were about an order of magnitude more potent in this assay than the best Fabs generated here). The class C Fabs interact with the internal coiled-coil of N-helices of gp41. Louis et al. [2005] (antibody binding site definition and exposure, antibody generation)

**No.** 780

MAb ID polyclonal

HXB2 Location gp160

Author Location gp120 (V3)

**Epitope** 

Subtype multiple

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Barin et al. 2005

Country France

Keywords acute/early infection, assay development

· A combination of 4 antigenic regions was used to differentiate between early (<180 days) and chronic infection. These regions were: p24; the gp41 peptide spanning the immunodominant epitope (IDE) of gp41, RVAVERYLKDQQLLGIWGC-SGKICTTAV, and a subtype D version of this peptide; 5 V3 consensus peptides including A, B, C, D, and CRF01-AE; and Integrase. V3 and the IDE provide the best discrimination, with >20 fold higher levels in chronic infection when assayed by EIA using dried serum spots. Antibodies to Integrase and p24 were not as distinctive, and people tend to lose, not increase, responses to p24 over time. This assay can be used to identify samples from early infection with high sensitivity and specificity. Barin et al. [2005] (assay development, acute/early infection)

**No.** 781

MAb ID polyclonal

HXB2 Location gp160

**Author Location** Env (gp160)

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Bettaieb et al. 1992

Keywords mimics

• gp160-specific Abs were detected in platelet elutates from HIV-1 infected patients with immunologic thrombocytopenia purpura (ITP). One patient with high titer anti-gp160/120 Abs had IgG that bound specifically to both gp160/120 and to platelet GPIIb/IIIa, apparent molecular mimicry. The crossreactive epitope on gp120 has not been defined, however a conformational aspect and/or glycosylation is likely to be involved. Bettaieb et al. [1992] (mimics)

No. 782

MAb ID 2F19C

HXB2 Location gp160

Author Location gp120 (HIV2ROD)

Epitope APGK

Neutralizing no

Immunogen vaccine

Vector/Type: peptide Strain: HIV-2 ROD

Species (Isotype) mouse

Ab Type gp120 C3

References Matsushita et al. 1995

• 2F19C: Binds in WB, but binds poorly to Env on the cell surface, APGK is the core binding region. Matsushita et al. [1995]

No. 783

**MAb ID** 1334-D (1334, 1334D)

HXB2 Location gp160

Author Location gp120 (HIV451)

**Epitope TRTSV** 

Neutralizing

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu) (NYU Med.

Center)

References Gorny et al. 2004; Gorny & Zolla-Pazner

2004; Nyambi et al. 2000; Gorny et al. 2000; Zolla-Pazner et al. 1999b; Zolla-Pazner et al.

1999a

Keywords antibody binding site definition and exposure

- 1334-D: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004]
- 1334-D: Called 1334. V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using HIV451 gp120. Gorny et al. [2004] (antibody

binding site definition and exposure)

• 1334-D: Called 1334 – binds to V3 peptides from MN, SF2, NY5, RF, and CDC4 strains as well as x-reactivity with peptides from A, C, D, F, G, and H subtypes - was suggested to be IgG11ambda here – binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared - no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and

Env Antibodies HIV Antibodies Tables

C5 tended to favor the monomer – V3 MAbs 447-52D, 838-D, and 1334 bound with a 7-10 fold preference for the oligomer. Gorny *et al.* [2000]

- 1334-D: Called 1334D A panel of 47 human MAbs was tested against 26 HIV-1 group M primary isolates from clades A through H 19 V3 MAbs were tested, and of 494 combinations, 44% displayed some viral binding V3 MAbs tended to have the most cross-reactive binding to clade A, B, C, and D isolates, less to E, F, G, and H 1334D showed intermediate cross-reactivity. Nyambi *et al.* [2000]
- 1334-D: This MAb was selected using oligomeric gp160 from HIV451. Zolla-Pazner *et al.* [1999a]
- 1334-D: MAb peptide-reactivity pattern clustered with immunological related MAbs: 1334, 419, 504, 447, 453 and 537 the core amino acids GP tended to be critical for reactivity in this group. Zolla-Pazner *et al.* [1999b]

## **IV-C-17** Env Antibodies

No. 784

MAb ID

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: gp120 Adjuvant: GM-CSF

Species (Isotype) mouse (IgG1)

References Rodríguez et al. 1999

• The murine Ab response to a chimeric of granulocyte-macrophage colony stimulating factor GM-CSF/gp120 in vaccinia was not higher titer that the response to a gp120-vaccinia construct, but the breadth of the Ab response was greater, in particular to the C-term region of gp120 – a cellular response of greater intensity was trigged to the GM-CSF/gp120 vaccinia construct, as measured by Elispot assay. Rodríguez *et al.* [1999]

**No.** 785

MAb ID

**HXB2 Location** Env

**Author Location** Env (384–467)

Epitope Neutralizing

Immunogen vaccine

Vector/Type: hepatitis B surface antigen lipoprotein particles (HsBAg) HIV compo-

nent: V3

Species (Isotype) macaque, rabbit

References Michel et al. 1993

• Immunization with recombinant HIV1 V3/HBsAg hybrid particles into rabbits or macaques elicited and maintained for several months anti-V3 or HIV-1 Env proliferative, CTL and Ab responses. Michel *et al.* [1993]

**No.** 786 **MAb ID** 

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing yes

Immunogen HIV-1 infection, vaccine

Species (Isotype) human

References Burton & Parren 2000

• This review article touches on why natural immune responses do not tend to favor potent neutralizing Ab production, and discusses possible vaccine strategies to counter this problem. Burton & Parren [2000]

No. 787

MAb ID

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Pellegrin et al. 1996

• Detection of an autologous NAb response in 12 patients with primary infections was delayed – for patients with a viral isolate obtained at month 1, autologous NAbs to viral isolates were generally not observed before month 6, and there was no apparent relationship between the emergence of neutralizing activity and the decrease of plasma viral load. Pellegrin *et al.* [1996]

No. 788

MAb ID

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype)

**References** Berger 2002

Keywords immunotherapy

• This medical hypothesis proposes that HIV shares domains with human proteins are masked from the immune response as they are seen as self. They propose blocking the shared determinants on human proteins in the thymus with antibodies, to allow anti-self responses which are normally inhibited to occur in HIV+ people. (immunotherapy)

**No.** 789

**MAb ID** 102-135

HXB2 Location Env

Author Location gp41 (HAM112, O group)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: O group

HAM112 HIV component: gp160

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Scheffel et al. 1999

**HIV Antibodies Tables Env Antibodies** 

Env were tested for MAb reactivity - 102-135 bound to two non-contiguous peptides in combination, assumed to form some type of helical structure, and not to either individually. Scheffel et al. [1999]

**No.** 790

**MAb ID** 1025

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

**Immunogen** 

Species (Isotype)

References Berman et al. 1997

• 1025: Binds to 1/7 isolates from breakthrough cases from a • 12H2: Env in a Semliki-Forest Virus (SFV) vector was used MN gp120 vaccine trial. Berman et al. [1997]

No. 791

MAb ID 105-134

**HXB2 Location** Env

**Author Location** gp41 (652–681 HAM112, O group)

**Epitope** Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: O group

HAM112 HIV component: gp160

**Species (Isotype)** mouse (IgG1 $\kappa$ )

References Scheffel et al. 1999

• 105-134: Overlapping peptides based on group O HAM112 Research Contact Evan Hersh and Yoh-Ichi Matsumoto Env were tested for MAb reactivity. Scheffel et al. [1999]

No. 792

**MAb ID** 10E9

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) mouse (IgG1)

References Papsidero et al. 1988

• 10E9: 100/100 HIV+ human sera could inhibit 10E9 binding. • 13.10: First HIV-1 specific human-mouse hybridoma that pro-Papsidero et al. [1988]

No. 793

MAb ID 126-50

**HXB2 Location** Env

Author Location gp41 (HXB2)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human ( $IgG2\kappa$ )

References Xu et al. 1991; Robinson et al. 1991; Tyler et al. 1990; Robinson et al. 1990b

- 126-50: No enhancing or neutralizing activity. Robinson et al.
- 126-50: Specific for a conformational epitope. Xu et al. [1991]
- 126-50: No enhancing activity for HIV-1 IIIB. Robinson et al. [1990b]

• 102-135: Overlapping peptides based on group O HAM112 • 126-50: Serves as target for antibody-dependent cellular cytotoxicity ADCC. Tyler et al. [1990]

No. 794

MAb ID 12H2

**HXB2 Location** Env

Author Location gp41 (530–677 HXB2)

**Epitope** 

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: Semliki-Forest Virus HIV com-

ponent: Env

**Species (Isotype)** mouse (IgM $\kappa$ )

References Giraud et al. 1999

to vaccinate mice intramuscularly as naked RNA, and an Ab response was induced to Env from which 12H2 was derived - and advantage of this method is that the protein is properly expressed. Giraud et al. [1999]

No. 795

**MAb ID** 13.10 (No. 13)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

References Wisnewski et al. 1996; Moran et al. 1993; Lake et al. 1989

• 13.10: NIH AIDS Research and Reference Reagent Program:

- 13.10: 13.10 is V H1 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski
- 13.10: Heavy (V HI) and light (V lambdaII) chain sequenced no enhancing or neutralizing activity - called No. 13. Moran et al. [1993]
- duces a MAb that binds to gp120 and gp160. Lake et al. [1989]

No. 796

MAb ID 1B1

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human

Research Contact Herman Katinger, Inst. Appl. Microbiol. Uni-

versity of Agricultural Science, Vienna, Aus-

References Kunert et al. 1998; Purtscher et al. 1994;

Buchacher et al. 1994

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• 1B1: The complete V, J and D(H) domain was sequenced – Species (Isotype) human (IgG1 $\kappa$ ) unlike non-neutralizing anti-gp41 MAb 3D6, five neutralizing MAbs (2F5, 2G12, 1B1, 1F7, and 3D5) showed extensive somatic mutations giving evidence of persistent antigenic pres- Research Contact Herman Katinger, Inst. Appl. Microbiol. or sure over long periods. Kunert et al. [1998]

• 1B1: Generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994]

**No.** 797

MAb ID 1D10

**HXB2 Location** Env

Author Location gp120 (34-55)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Species (Isotype)

Research Contact Phil Berman

References Callahan et al. 1991

- · Isolation of antibody.
- 1D10: Polyanionic polysaccharides were proposed to inhibit viral functions such as binding and syncytia formation through interations mediated through the local high positive charge density in the V3 loop. The binding of this N-term binding antibody is increased by dextransulfate, in contrast to anti-V3 antibodies that are inhibited. Callahan et al. [1991]

No. 798

MAb ID 1F7

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human

Research Contact Herman Katinger, Inst. Appl. Microbiol. University of Agricultural Science, Vienna, Aus-

References Grant et al. 2000; Kunert et al. 1998; Purtscher et al. 1994; Buchacher et al. 1994

- 1F7: There is an anti-idiotype MAb named 1F7 that was raised against pooled IgG from HIV-1 + subjects that recognizes a set of antibodies against HIV Gag, Pol, and Env, and this MAb is reported to inhibit anti-HIV CTL activity—this is not the same as the 1F7 described by Buchacher et al.. Grant et al. [2000]
- 1F7: The complete V, J and D(H) domain was sequenced unlike non-neutralizing anti-gp41 MAb 3D6, five neutralizing MAbs (2F5, 2G12, 1B1, 1F7, and 3D5) showed extensive somatic mutations giving evidence of persistent antigenic pressure over long periods. Kunert et al. [1998]
- 1F7: Generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher et al. [1994]

No. 799

MAb ID 2G12 (c2G12)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Subtype B

**Neutralizing** LP

Immunogen HIV-1 infection

**Ab Type** gp120 carbohydrates at glycosylation residues in C2, C3, C4, and V4

Polymun Scientific Inc., Vienna, Austria,

References Selvarajah et al. 2005; Raviv et al. 2005; Pinter et al. 2005; Pashov et al. 2005; Nakowitsch et al. 2005; Martín-García et al. 2005; Lusso et al. 2005; Louis et al. 2005; Krachmarov et al. 2005; Haynes et al. 2005; Gorny et al. 2005; Chen et al. 2005; Calarese et al. 2005; Wang et al. 2004; Safrit et al. 2004; Pugach et al. 2004; Pinter et al. 2004; Pantophlet et al. 2004; Opalka et al. 2004; Nabatov et al. 2004; Lorin et al. 2004; Liao et al. 2004; Jeffs et al. 2004; Ferrantelli et al. 2004a; Ferrantelli et al. 2004b; Dacheux et al. 2004; Biorn et al. 2004; Binley et al. 2004; Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Wolbank et al. 2003; Ohagen et al. 2003; Montefiori et al. 2003; Louis et al. 2003; Kitabwalla et al. 2003; Raja et al. 2003; Singh et al. 2003; Wang 2003; Richman et al. 2003; Mascola 2003; Mascola et al. 2003; Hart et al. 2003; Ferrantelli et al. 2003; Dey et al. 2003; Cavacini et al. 2003; Binley et al. 2003; Abrahamyan et al. 2003; Albu et al. 2003; Herrera et al. 2003; Pantophlet et al. 2003a; Calarese et al. 2003; Stiegler et al. 2002; Kwong et al. 2002; Gorry et al. 2002; Cavacini et al. 2002; Bures et al. 2002; Liu et al. 2002; Ferrantelli & Ruprecht 2002; Zhang et al. 2002; Mascola 2002; Grundner et al. 2002; Edwards et al. 2002; Armbruster et al. 2002; Chakrabarti et al. 2002; Xu et al. 2002; Yang et al. 2002; Schulke et al. 2002; Scanlan et al. 2002; Sanders et al. 2002; Golding et al. 2002b; Savarino et al. 2001; Xu et al. 2001; Hofmann-Lehmann et al. 2001; Spenlehauer et al. 2001; Stiegler et al. 2001; Verrier et al. 2001; Zeder-Lutz et al. 2001; Poignard et al. 2001; Moore et al. 2001; Barnett et al. 2001; Zwick et al. 2001c; Mascola & Nabel 2001; Si et al. 2001; Park et al. 2000; Grovit-Ferbas et al. 2000; Baba et al. 2000; Robert-Guroff 2000; Binley et al. 1999; Mascola et al. 2000; Mascola et al. 1999; Parren et al. 1999; Poignard et al. 1999; Crawford et al. 1999; Altmeyer et al. 1999; Beddows et al. 1999; Montefiori & Evans 1999; Schonning et al. 1998; Kunert et al. 1998; Frankel et al. 1998; Wyatt & Sodroski 1998; Li et al. 1998; Parren et al. 1998b; Takefman et al. 1998; Fouts et al. 1998; Trkola et al. 1998; Binley et al. 1998; Connor et al. 1998; Sullivan et al. 1998b; Parren et al. 1998a; Mondor et al. 1998; Wyatt et al. 1998; Andrus et al. 1998; Parren et al. 1997b; Burton & Montefiori 1997; Ugolini et al. 1997; Mas**HIV Antibodies Tables Env Antibodies** 

> cola et al. 1997; Moore & Trkola 1997; Li et al. 1997; Fouts et al. 1997; Binley et al. 1997a; Mo et al. 1997; D'Souza et al. 1997; Sattentau 1996; Trkola et al. 1996a; Poignard et al. 1996b; Moore & Sodroski 1996; Trkola et al. 1996b; McKeating 1996; McKeating et al. 1996; Moore & Ho 1995; Trkola et al. 1995; Buchacher et al. 1994

Keywords acute/early infection, antibody binding site definition and exposure, antibody generation, antibody interactions, antibody sequence, variable domain, assay development, assay standardization/improvement, autologous responses, brain/CSF, co-receptor, complement, escape, immunoprophylaxis, immunotherapy, isotype switch, kinetics, mother-to-infant transmission, mucosal immunity, neutralization potency, reversion, viral fitness, review, structure, subtype comparisons, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or crossneutralization

- 2G12: UK Medical Research council AIDS reagent: ARP3030.
- 2G12: NIH AIDS Research and Reference Reagent Program: 1476.
- 2G12: Precise characterization of 2G12 binding to carbohydrate was undertaken; the 2G12 Fab was co-crystallized with four oligomannose derivatives, Man4, Man5, Man7 and Man8. 2G12 recognizes the terminal Manα1-2Man both in the context of the D1 arm (Man $\alpha$ 1-2Man $\alpha$ 1-2Man) and D3 arm (Man $\alpha$ 1-2Manα1-6Man) of the Man9GlcNAc2 moiety, but not the D2 arm. This gives the 2G12 more binding flexibility than previously thought, as only the D1 arm binding had been shown previously. Calarese et al. [2005] (antibody binding site definition and exposure, structure)
- 2G12: The lack of glycosylation sites at residues Asn 295 and Thy 394 within C-clade gp120s generally causes the loss of 2G12 recognition. Introduction of glycans in the subtype C strain HIV-1CN54 at these positions restored 2G12 binding, and addition of just a single glycan partially restored binding (V295N + A394T » V295N > A395T). 2G12 epitope recovery decreased b12 binding. Chen et al. [2005] (antibody binding site definition and exposure)
- 2G12: 2909 is a human anti-Env NAb that was selected by a neutralization assay and binds to the quaternary structure on the intact virion. ELISA-based competition assays and subsequent mutational analysis determined that the CD4BS and V2 and V3 loops contribute to the 2909 epitope: 2909 binding was inhibited by MAbs 447-52d (anti-V3), 830A (anti-V2), and IgG1b12 (anti-CD4BS) and sCD4. 2909 was not inhibited by MAbs 670, 1418, nor 2G12; in fact, 2G12 enhanced 2909 binding. Gorny et al. [2005]
- 2G12: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. Unlike the other three broadly neutralizing human anti-HIV-1 MAbs, 2G12 has no indication of polyspe-

cific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

- 2G12: Sera from subtype A infected individuals from Cameroon have antibodies that react strongly with subtype A and subtype B V3 loops in fusion proteins, and neutralize SF162 pseudotypes, while sera from 47 subtype B infected individuals reacted only with subtype B V3s. Sera from Cameroon did not neutralize primary A or B isolates, due to indirect masking by the V1/V2 domain rather than due to loss of the target epitope. Neutralization by Cameroonian sera MAbs was blocked by Clade A and B V3 loop fusion proteins, while NAbs to non-V3 epitopes, 2F5, 2G12, and b12, were not blocked. Krachmarov et al. [2005] (antibody binding site definition and exposure, variant cross-recognition or crossneutralization, subtype comparisons)
- 2G12: Nine anti-gp41 bivalent Fabs that interacted with either or both of the six-helix bundle and the internal coiled-coil of N-helices of gp41 were selected from a non-immune human phage display library. The IC50 the range for the inhibition of LAV ENV-mediated cell-fusion was 6-61 ug/ml – for context, 2F5 and 2G12 (IC50s of 0.5-1.5 ug/ml) were about an order of magnitude more potent in this assay than the best Fabs generated here. Louis *et al.* [2005] (**neutralization potency**)
- 2G12: The HIV-1 Bori-15 variant was adapted from the Bori isolate for replication in microglial cells. Bori-15 had increased replication in microglial cells and a robust syncytium-forming phenotype, ability to use low levels of CD4 for infection, and increased sensitivity to neutralization by sCD4 and 17b. Four amino acid changes in gp120 V1-V2 were responsible for this change. Protein functionality and integrity of soluble, monomeric gp120-molecules derived from parental HIV-1 Bori and microglia-adapted HIV-1 Bori-15 was assessed in ELISA binding assays using CD4BS MAbs F105 and IgG1b12, glycanspecific 2G12, and V3-specific 447-52D, and were unchanged. Association rates of sCD4 and 17b were not changed, but dissociation rates were 3-fold slower for sCD4 and 14-fold slower for 17b. Martín-García et al. [2005] (antibody binding site definition and exposure)
- 2G12: Passive immunization of 8 HIV-1 infected patients with 4E10, 2F5 and 2G12 (day 0, 4E10; days 7, 14 and 21 4E10+2G12+2F5; virus isolated on days 0 and 77) resulted in 0/8 patients with virus that escaped all three NAbs. Three patients had viruses that escaped 2G12, and two of these were sequenced. Each had lost two of the glycosylation sites required for 2G12 binding (one had 295 N->D and 332 N->T changes, the other had 295 N->T and 392 N->T changes). In a companion in vitro study, resistance to a single MAb emerged in 3-22 weeks, but triple combination resistance was slower and characterized by decreased viral fitness. In contrast to the in vivo escape study, only one N was lost in the in vitro experiments, a 386 N->K change in a triple resistant mutant. The lack of resistance to the combination of MAbs in vivo and the reduced fitness of the escape mutants selected in vitro suggests passive immunotherapy may be of value in HIV infection. Nakowitsch et al. [2005] (escape, immunotherapy)
- 2G12: Concanavalin A (ConA) binds to mannose and blocks 2G12 binding, but 2G12 does not block ConA binding. ConA binding is less sensitive to mutations in glycosylation sites than 2G12. Furthermore, ConA neutralizes HIV-1 at a post-CD4

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binding step. Thus, this report indicates that designing antigens based on the HIV-1 mannose residues that bind ConA may be an effective vaccine strategy, as antibodies elicited might be broadly cross-reactive. Pashov *et al.* [2005] (vaccine antigen design)

- 2G12: This study is about the V2 MAb C108g, that is type-specific and neutralizes BaL and HXB2. JR-FL is a neutralization resistant strain; modification of JRFL at V2 positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The modification in V2 also increased neutralization sensitivity to V3 MABs 4117c, 2219, 2191, and 447-52D, but only had minor effects on neutralization by CD4BS MAb 5145A, and broadly neutralizing MAbs IgG1b12, 2G12, and 2F5. Pinter et al. [2005] (antibody binding site definition and exposure)
- 2G12: Retrovirus inactivation for vaccine antigen delivery was explored through lipid modification by hydrophobic photoin-duced alkylating probe 1.5 iodonaphthylazide (INA). The viral proteins were shown to be structurally intact in the treated non-infectious virus, through the preservation of antibody binding sites for polyclonal anti-gp120 serum, and for broadly neutralizing MAbs 2G12, b12 and 4E10, although the modifications of the lipid disabled viral infection. Raviv et al. [2005] (vaccine antigen design)
- 2G12: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. 2G12 had diminished binding to both antigen constructs. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)
- 2G12: 93 viruses from different clades were tested for their neutralization cross-reactivity using a panel of HIV antibodies. 2G12 primarily neutralized B clade viruses with sporadic neutralization of A, D, and two AC recombinants, and no C or CRF01 (E) isolates. Envelopes from subtypes C and E have generally lost critical glycans for 2G12 binding. Binley *et al.* [2004] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 2G12: The peptide 12p1 (RINNIPWSEAMM) inhibits direct binding of YU2 gp120 or Env trimer to CD4, CCR5 and MAb 17b in a concentration-dependent allosteric manner. 12p1 is thought to bind to unbound gp120 near the CD4 binding site, with a 1:1 stoichiometry. 12p1 also inhibited MAb F105 binding. presumably because F105 favors an unactivated conformation, but not MAbs 2G12 or b12. The 1:1 stoichiomtry, the fact that the peptide binding site is accessible on the trimer, the non-CD4 like aspect of the binding, and an ability to inhibit viral infection in cell cultures make it a promising lead for therapeutic design. Biorn *et al.* [2004]
- 2G12: Env sequences were derived from 4 men at primary infection and four years later; the antigenicity in terms of the ability to bind to 2G12, 2F5 and IgG1b12 was determined. 2G12 bound primarily to late clones in 3 of the 4 patients, and to both early and late in the other patient. Neither 2F5 nor

- IgG1b12 showed a difference in binding affinity to early or late envelopes. The number of glycosylation sites increased in the three patients. The ability to bind to 2G12 correlated perfectly with having all three sites known to be important for binding: N295 in C2, N332 in C3, and N392 in the V4 loop. Dacheux *et al.* [2004] (antibody binding site definition and exposure, acute/early infection, kinetics)
- 2G12: Neonatal rhesus macaques were exposed orally to a pathogenic SHIV, 89.6P. 4/8 were given an intramuscular, passive immunization consisting of NAbs 2G12, 2F5 and 4E10, each given at a different body sites at 40 mg/kg per Ab, at one hour and again at 8 days after exposure to 89.6P. The four animals that were untreated all died with a mean survival time of 5.5 weeks, the four animals that got the NAb combination were protected from infection. This model suggests Abs may be protective against mother-to-infant transmission of HIV. Ferrantelli *et al.* [2004b] (mother-to-infant transmission)
- 2G12: Nabs against HIV-1 M group isolates were tested for their ability to neutralize 6 randomly selected HIV-1 O group strains. 2G12 didn't neutralizeO group strains, although it was included in a quadruple combination of b12, 2F5, 2G12, and 4E10, that neutralized the six Group O viruses between 62-97%. Ferrantelli *et al.* [2004a] (variant cross-recognition or cross-neutralization)
- 2G12: This paper is a reviw of anti-HIV-1 Envelope antibodies.
   This unique epitope is formed from carbohydrates. The mechanism of MAb neutralization is thought to be steric inhibition of CCR5 binding. 2G12 neutralizes many TCLA strains and about 40% of primary isolates tested. Gorny & Zolla-Pazner [2004] (review)
- 2G12: A set of oligomeric envelope proteins were made from six primary isolates for potenial use as vaccine antigens: 92/UG/037 (clade A), HAN2/2 (clade B), 92/BR25/025 (clade C), 92/UG/021 (clade D), 93/BR/029 (clade F) and MVP5180 (clade O). This was one of a panel of MAbs used to explore folding and exposure of well characterized epitopes. The clade C isolate BR25 is apparently misfolded, as conformation-dependent antibodies did not bind to it. 2G12 bound to clade A, B, D and F HIV-1 primary isolates. Polyclonal sera raised in rabbits against these antigens cross-bound the other antigens, but none of the sera had neutralizing activity. Jeffs *et al.* [2004] (vaccine antigen design, subtype comparisons)
- 2G12: 2G12 was used as a positive control in a study that showed that A32-rgp120 complexes open up the CCR5 coreceptor binding site, but did not induce neutralizing antibodies with greater breadth among B subtype isolates than did uncomplexed rgp120 in vaccinated guinea pigs. Liao et al. [2004] (vaccine antigen design)
- 2G12: Mice susceptible to MV infection were intraperitoneally immunized with native HIV-1 89.6 env gp160 and gp140 and δV3 HIV-1 89.6 mutants expressed in live attenuated Schwarz measles vector (MV). The gp160ΔV3 construct raised more cross-reactive NAbs to primary isolates. A HIVIG/2F5/2G12 combination was used as a positive control and could neutralize all isolates. Lorin *et al.* [2004] (vaccine antigen design)
- 2G12: A set of HIV-1 chimeras that altered V3 net charge and glycosylation patterns in V1V2 and V3, involving inserting V1V2 loops from a late stage primary isolate taken after the R5 to X4 switch, were studied with regard to phenotype, co-

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receptor usage, and MAb neutralization. The loops were cloned into a HXB2 envelope with a LAI viral backbone. It was observed that the addition of the late-stage isolate V1V2 region and the loss of V3-linked glycosylation site in the context of high positive charge gave an X4 phenotype. R5X4 viruses were more sCD4 and 2G12 neutralization resistant than either R5 or X4, but the opposite pattern was observed for b12. Addition of the late stage V1V2 altered neutralization for both MAbs, but this alteration was reversed with the loss of the V3 glycan. Nabatov *et al.* [2004] (antibody binding site definition and exposure, co-receptor)

- 2G12: An antigen panel representing different regions of gp41 was generated, and sera from 23 individuals were screened.
   2G12 was a control, binding to gp120 but to none of the gp41 peptides in the experiment. Opalka *et al.* [2004] (assay development, assay standardization/improvement)
- 2G12: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it binds to the neutralizing MAb 2G12. It masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120. Pantophlet *et al.* [2004] (vaccine antigen design)
- 2G12: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. 2G12 was the only MAb that neutralized JRFL more efficiently than SF162, with a 6-fold lower ND50 for JRFL. 2G12 also had ahigher affinity for JRFL. Pinter et al. [2004] (variant cross-recognition or cross-neutralization)
- 2G12: A primary isolate, CC1/85, was passaged 19 times in PBMC and gradually acquired increased sensitivity to FAb b12 and sCD4 that was attributed to changes in the V1V2 loop region, in particular the loss of a potential glycosylation site. The affinity for sCD4 was unchanged in the monomer, suggesting that the structural impact of the change was manifested at the level of the trimer. The passaged virus, CCcon19, retained an R5 phenotype and its neutralization susceptibility to other Abs was essentially the same as CC1/85. The IC50 for 2G12 was 1.8 for CC1/85, and was 4.2 for CCcon19, so both the primary and passaged viruses were neutralized. Pugach *et al.* [2004] (reversion, viral fitness, variant cross-recognition or cross-neutralization)
- 2G12: This review discusses research presented at the Ghent Workshop of prevention of breast milk transmission and immunoprophylaxis for HIV-1 in pediatrics (Seattle, Oct. 2002), and makes the case for developing passive or active immunoprophylaxis in neonates to prevent mother-to-infant transmission. Macaque studies have shown that passive transfer of

NAb combinations (for example, IgG1b12, 2G12, 2F5, and 4E10; or 2G12 and 2F5) can confer partial or complete protection to infant macaques from subsequent oral SHIV challenge. Safrit *et al.* [2004] (**immunoprophylaxis**, **mother-to-infant transmission**)

- 2G12: Synthetic mannose Man9 clusters arranged on a scaffold were used to mimic the epitope of 2G12. Bi-, tri, and tetra-valent clusters had a 7-, 22-, and 73-fold higher affinities for 2G12 than the monomers, suggesting that 2G12 binds best to multiple carbohydrate moieties. 2G12 bound larger mannose oligosaccharides with higher affinity: Ma9GlcNAc bound 210- and 74-fold more effectively that Man6GlcNac and Man5GlcNAc, respectively. Wang et al. [2004] (antibody binding site definition and exposure)
- 2G12: SOS-Env is a mutant protein engineered to have a disulfid bond between gp120 and gp41. Cells expressing SOS-Env due not fuse with target cells expressing CD4 and CCR5, although the fusion process proceeds to an intermediate state associated with CD4 and co-receptors, prior to the formation of the six helix bundle that allows fusion.2G12 was used to monitor surface expression of SOS-Env compared to wildtype. Abrahamyan *et al.* [2003] (**co-receptor**, **vaccine antigen design**)
- 2G12: 2G12 was used as a positive control to test for a NAb activity in mice intranasally immunized with gp120 or gp140 with IL-12 and Cholera Toxin B. Albu et al. [2003]
- 2G12: The SOS mutant envelope protein introduces a covalent disulfide bond between gp120 surface and gp41 transmembrane proteins into the R5 isolate JR-FL by adding cysteines at residues 501 and 605. Pseudovirions bearing this protein bind to CD4 and co-receptor bearing cells, but do not fuse until treatment with a reducing agent, and are arrested prior to fusion after CD4 and co-receptor engagement. 2G12 is able to neutralize both the wildtype and SOS protein comparably, but 2G12 could not neutralize SOS when added post-attachment. Binley *et al.* [2003] (vaccine antigen design)
- 2G12: Crystal structure analysis of Fab 2G12 alone or complexed with Manα1-2Man or Man9GlcNac2 demonstrates that the exchange of VH domains forms stable dimers for gp120 binding. Two Fabs assemble in an interlocked VH domain swapped dimer, providing an extended surface for multivalent interaction with the cluster of oligomannose on gp120, allowing high-affinity recognition of repeated epitopes in the carbohydrate structure. Ala substitutions of the 2G12 VH/VH' interface residues Ile H19, Arg H57, Phe H77, Tyr H80, Val H84 and Pro H113 result in the loss of 2G12-gp120 JR-FL binding. Calarese *et al.* [2003] (antibody binding site definition and exposure, antibody sequence, variable domain, structure)
- 2G12:The MAb B4e8 binds to the base of the V3 loop, neutralizes multiple primary isolates and was studied for interaction with other MAbs. B4e8 and 2G12 enhanced each others binding, and gave synergistic neutralization. B4e8 could neutralize R5X4 virus 92HT593 better than 2G12, while 2G12 was better at neutralizing R5 virus 92US660. Cavacini et al. [2003] (antibody interactions)
- 2G12: A sCD4-17b single chain chimera was made that can bind to the CD4 binding site, then bind and block co-receptor interaction. This chimeric protein is a very potent neutralizing

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agent, more potent than IgG1b12, 2G12 or 2F5 against Ba-L infection of CCR5-MAGI cells. It has potential for prophylaxis or therapy. Dey *et al.* [2003] (**co-receptor**)

- 2G12: Four newborn macaques were challenged with pathogenic SHIV 89.6 and given post exposure prophylaxis using a combination of NAbs 2F5, 2G12, 4E10 and IgG1b12. 2/4 treated animals did not show signs of infection, and 2/4 macaques maintained normal CD4+ T cell counts and had a lower delayed peak viremia compared to the controls. Ferrantelli *et al.* [2003] (immunoprophylaxis, mother-to-infant transmission)
- 2G12: This study investigates the effects of glycosylation inhibitors on the binding between HIV-1 gp120 and mannose-binding lectin (MBL). Mannosidase I inhibitor deoxymanno-jirimycin (dMM) inhibits formation of complex and hybrid N-linked saccharides and yields virus with more mannose residues. dMM added during viral production significantly enhanced the binding 2F5 and 2G12, but not IgG1b12 in a viral capture assay. Hart et al. [2003] (antibody binding site definition and exposure)
- 2G12: CD4BS MAbs b12 (neutralizing) and 205-42-15, 204-43-1, 205-46-9 (non-neutralizing) all cross-competed for binding to monomeric gp120, indicating the topological proximity of their epitopes, however, the non-neutralizing CD4BS MAbs did not interfere with the neutralization activity of MAb b12 2G12 was used to normalize and as a control in these experiments. Herrera et al. [2003] (antibody interactions)
- 2G12: MAbs IgG1b12, 2G12, 2F5 and 4E10 were tested for their ability to neutralize two primary HIV-1 clade A isolates (UG/92/031 and UG/92/037) and two primary HIV-1 clade D isolates (UG/92/001 and UG/92/005). 4E10 demonstrated the most potent cross-neutralization activity. Quadruple administration of MAbs IgG1b12, 2G12, 2F5, and 4E10 induced strong synergistic neutralization of 4 clade A isolates (UG/92/031, UG/92/037, RW/92/020 and RW/92/025) as well as 5 clade D isolates (UG/92/001,UG/9/005, /93/086/RUG/94/108, UG/94/114). The authors note this combination of 4 MAbs neutralizes primary HIV A, B, C, and D isolates. Kitabwalla et al. [2003] (antibody interactions, immunoprophylaxis, variant cross-recognition or cross-neutralization, mother-to-infant transmission, subtype comparisons)
- 2G12: Polyclonal Abs raised against soluble trivalently linked N35CCG-N13 and N34CCG, the internal trimeric core of the coiled-coil ectodomain, inhibit HIV-1 Env-mediated cell fusion at levels comparable to 2G12. Louis *et al.* [2003] (vaccine antigen design)
- 2G12: This review dicusses the importance and function of protective antibody responses in animal model studies in the context of effective vaccine development. SHIV models have shown protection using high levels of MAbs can prevent infection, and partial protection that can influence disease course can be obtained from modest levels of NAbs. SHIV challenges studies conducted with infusions of combinations of MAbs b12, 2G12, and 2F5 are reviewed. Mascola [2003] (immunoprophylaxis, review)
- 2G12: Infusions of 2F5 and 2G12 intravenously administered 24h prior to vaginal SHIV-89.P challenge are able to protect macaques from infections. Animals that recieve a IL-2 adjuvanted DNA immunization SIV Gag and HIV Env have T-cell

- responses and lower viral loads, but were not protected. Suboptimal levels of 2F5 and 2G12 were not able to confer sterile protection in combination with the T-cell responses stimulated by DNA immunizations. Mascola *et al.* [2003]
- 2G12: AC10 is a subject who was given treatment early after infection, and had a viral rebound after cessation of therapy, which then declined to a low level. The polyclonal sera from AC10 could potently neutralize the rebound virus, and NAb escape followed with a neutralizing response against the escape variant and subsequent escape from that response. Viral loads remained low in this subject despite escape. The rebound isolate that was potently neutralized by autologous sera was not particularly neutralization sensitive, as it resisted neutralization by sCD4 and MAbs IgG1b12, 2G12 and 2F5, and was only moderately sensitive to sera from other HIV+ individuals that had high titers of NAbs to TCLA strains. Montefiori *et al.* [2003] (acute/early infection, escape)
- 2G12: Env genes derived from uncultured brain biopsy samples from four HIV-1 infected patients with late-stage AIDS were compared to env genes from PBMC samples. Brain isolates did not differ in the total number or positions of N-glycosylation sites, patterns of coreceptor usage, or ability to be recognized by gp160 and gp41 MAbs. 2G12 was the only MAb tested to recognize all blood and brain isolates from all four patients by gp120 immunoprecipitation. Ohagen *et al.* [2003] (variant cross-recognition or cross-neutralization)
- 2G12: Alanine scanning mutagenesis was used to compare substitutions that affected anti-CD4BS NAb b12 rec gp120s were engineered to contain combinations of Alanine substitutions that enhanced b12 binding, and while binding of b12 to these gp120 monomers was generally maintained or increased, binding by five non-neutralizing anti-CD4bs MAbs (b3, b6, F105, 15e, and F91) was reduced or completely abolished 2G12 binding was largely unperturbed, indicating these proteins were not grossly misfolded. Pantophlet *et al.* [2003a] (antibody binding site definition and exposure)
- 2G12: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of non-neutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet *et al.* [2003b] (vaccine antigen design)
- 2G12: This paper shows that binding of CD4BS MAbs to Env blocks the conformational shift that allows co-receptor CCR5 binding and CD4-independent mediated cell fusion. CD4BS MAbs IgG1b12, F91 and F105 and their Fab counterparts (except for C11, used as a negative control) inhibited CD4-independent JR-FL and YU-2 gp120-CCR5 binding to CCR5-expressing Cf2Th cells and syncytium formation. The carbohydrated binding MAb 2G12 also inhibited CD4-independent syncytium formation. Raja et al. [2003] (co-receptor)
- 2G12: Most plasma samples of patients from early infection had NAb responses to early autologous viruses, and NAbs against heterologous strains tended to be delayed. Serial plasma samples were tested against serial isolates, and neutralization escape was shown to be rapid and continuous throughout infection. Autologous neutralization-susceptible and resistant

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viruses from four patients were tested for susceptability to neu• 2G12: This study examined Ab interactions, binding and neutralizing Ab responses using MAbs 2G12, IgG1b12 and 2F5. No correlation was established, all viruses tested were susceptible to at least one of the neutralizing MAbs. Two patients that did not have an autologous NAb response also did not evolve changes in susceptibility to these MAbs, while one patient with a pattern of autologous neutralization and escape acquired a 2G12 sensitive virus at month 6, and lost IgG1b12 sensitivity at month 21. Richman et al. [2003] (autologous responses, acute/early infection, escape)

- 2G12: To begin to design vaccine antigens that can mimic the carbohydrate structure, the gp120 peptide 336-342 was synthesized with Man(9), Man(6), and Man(5) moieties attached. Singh et al. [2003] (vaccine antigen design)
- 2G12: Review of current neutralizing antibody-based HIV vaccine candidates and strategies of vaccine design. Strategies for targeting of the epitopes for NAbs 2F5, 2G12, 4E10, b12, and Z13 are described. They have shown that both N-glycans, at 295N and 332N are required for 2G12 binding, emphasizing the oligosaccharide cluster nature of the epitope, and suggest the uniqueness of the target structure may not result in autoimmune reactions. Wang [2003] (vaccine antigen design, review)
- 2G12: The broadly neutralizing antibodies 2F5 and 2G12 were class-switched from IgG to IgA and IgM isotypes. Neutralizing potency was increased with valence for 2G12 so the IgM form was most potent, but for 2F5 the IgG form was most potent. Eight primary isolates were tested including two subtype A isolates. The polymeric IgM and IgA Abs, but not the corresponding IgGs, could interfere with HIV-1 entry across a mucosal epithelial layer, although they were limited in a standard neutralization assay. All isotypes could interact with activated human sera, presumably through complement, to inhibit HIV replication. Wolbank et al. [2003] (complement, isotype switch, variant cross-recognition or cross-neutralization, mucosal immunity, subtype comparisons)
- 2G12: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. 2G12 had no impact on 4KG5 binding. Zwick et al. [2003] (antibody interactions)
- 2G12: A phase I trial in seven HIV+ individuals was conducted with MAbs 2F5 and 2G12 - no clinical or laboratory abnormalities were observed throughout the study – eight infusions were administered over a 4-week period (total dose 14 g) - the elimination half-life  $(t_1/2)$  was calculated to be 7.94 (range, 3.46–8.31) days for 2F5 and 16.48 (range, 12.84–24.85) days for 2G12. Armbruster et al. [2002] (kinetics, immunother-
- 2G12: IgG1b12 neutralized many South African (5/8) and Malawian (4/8) clade C primary HIV-1 isolates, being more effective than 2F5 which neutralized only two Malawian and no South African isolates. 2G12 did not neutralize any of the 16 isolates. Bures et al. [2002] (subtype comparisons)

- tralization with a B clade R5 isolate (92US660) and R5X4 isolate (92HT593). Abs generally bound and neutralized the R5X4 isolate better than the R5 isolate. Anti-gp41 MAb F240 did not affect binding of 2G12 to either R5X4 and R5 isolates, and anti-V3 MAb B4a1 increased 2G12 binding to R5X4 virions but not R5. Neutralization with B4al and 2G12 was additive for the R5X4 virus, and was enhanced for the R5 virus. Cavacini et al. [2002] (antibody interactions, co-receptor, variant cross-recognition or cross-neutralization)
- 2G12: A modified gp140 (gp140deltaCFI), with C-term mutations intended to mimic a fusion intermediate and stabilize trimer formation, retained antigenic conformational determinants as defined by binding to CD4 and to MAbs 2F5, 2G12, F105, and b12, and enhanced humoral immunity without diminishing the CTL response in mice injected with a DNA vaccine. Chakrabarti et al. [2002] (vaccine antigen design)
- 2G12: Truncation of the gp41 cytoplasmic domain of X4, R5, and X4R5 viruses forces a conformation that more closely resembles the CD4 bound state of the external Envelope, enhancing binding of CD4i MAbs 17b and 48d and of CD4BS MAbs F105, b12, and in most cases of glycosylation site dependent MAb 2G12 and the anti-gp41 MAb 246D - in contrast, binding of the anti-V2 MAb 697D and the anti-V3 MAb 694/98D were not affected - viruses bearing the truncation were more sensitive to neutralization by MAbs 48d, b12, and 2G12 - the anti-C5 MAb 1331A was used to track levels of cell surface expression of the mutated proteins. Edwards et al. [2002] (antibody binding site definition and exposure)
- 2G12: Review of NAbs that notes 2G12 alone or in combination with other MAbs can protect some macaques against SHIV infection, that it has strong ADCC activity, and that it is safe and well tolerated in humans. Ferrantelli & Ruprecht [2002] (immunoprophylaxis)
- 2G12: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry - preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion – the preincubation 31.5 C step did not alter the inhibitory activity of neutralizing Abs anti-gp41 2F5, or anti-gp120 2G12, IG1b12, 48d, and 17b. Golding et al. [2002b] (antibody binding site definition and exposure)
- 2G12: UK1-br and MACS2-br are R5 isolates derived from brain tissue samples from AIDS patients with dementia and HIV-1 encephalitis; both are neurotropic, but only UK1-br induced neuronal apoptosis and high levels of syncytium formation in macrophages. UK1-br Env had a greater affinity for CCR5 than MACS-br, and required low levels of CCR5 and CD4 for cell-to-cell fusion and single round infection. PBMC infected with UK1-br and MACS2-br virus isolates were resistant to neutralization by MAb 2G12. UK1-br was more sensitive than MACS2-br to IgG1b12, 2F5 and CD4-IgG2 neutralization. Gorry et al. [2002] (brain/CSF, co-receptor)
- 2G12: HIV-1 gp160deltaCT (cytoplasmic tail-deleted) proteoliposomes (PLs) containing native, trimeric envelope glycoproteins from R5 strains YU2 and JRFL, and X4 strain HXBc2, were made in a physiologic membrane setting as candidate immunogens for HIV vaccines - 2F5 bound to gp160deltaCT

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with a reconstituted membrane ten-fold better than the same protein on beads, while such an affinity difference was not seen with F105 and 2G12 – anti-CD4BS MAbs IgG1b12 and F105, A32 (C1-C4), C11 (C1-C5), and 39F (V3) MAbs bound gp160deltaCT PLs indistinguishably from gp160deltaCT expressed on the cell surface. Grundner *et al.* [2002] (antibody binding site definition and exposure, vaccine antigen design)

- 2G12: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, except for 2G12, which might not have bound well to the carbohydrate additions on the Drosophila expressed core. Enthalpy and entropy changes were divergent, but compensated. Not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs (17b, 48d, 1.5e, b6, F105 and F91) had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, but the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. The high values suggest surface burial or protein folding an ordering of amino acids. 2G12 had an entropy value of -1.6. These results suggest that while the trimeric Env complex has four surfaces, a nonneutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong et al. [2002] (antibody binding site definition and exposure)
- 2G12: Review of NAbs that discusses mechanisms of neutralization, passive transfer of NAbs and protection in animal studies, and vaccine strategies. Liu *et al.* [2002] (**review**)
- 2G12: Rhesus macaques were better protected from vaginal challenge with SHIV89.6D (MAb 2G12, 2/4; MAbs 2F5/2G12, 2/5; and HIVIG/2F5/2G12, 4/5 infected) than from intravenous challenge (MAb 2G12, 0/3; MAbs 2F5/2G12, 1/3; and HIVIG/2F5/2G12, 3/6 infected)—the animals that were infected by vaginal challenge after Ab infusion had low or undetectable viral RNA levels and modest CD4 T-cell decline. Mascola [2002] (immunoprophylaxis, mucosal immunity)
- 2G12: The 2G12 epitope is composed of carbohydrates involving high-mannose and hybrid glycans of residues 295, 332, and 392, with peripheral glycans from 386 and 448 contributing on either flank, and with little direct gp120 protein surface involvement these mannose residues are proximal to each other near the chemokine receptor binding surface. Sanders et al. [2002] (antibody binding site definition and exposure)
- 2G12: Alanine scanning mutagenesis used in conjunction with competition and replacement studies of N-linked carbohydrates and sugars suggest that the 2G12 epitope is formed from mannose residues contributed by the glycans attached to N295 and N332, with the other N-linked carbohydrates in positions N339, N386, and N392 playing a role in maintaining conformation relevant to 2G12 binding – N295A and N332A mutants showed essentially unchanged anti-CD4BS NAb b12 binding affinities, while N339A, N386A and N392A mutants displayed

significantly lowered b12 affinity, presumably due to conformational changes. Scanlan *et al.* [2002] (**antibody binding site definition and exposure**)

- 2G12: Ab binding characteristics of SOS gp140 were tested using SPR and RIPA SOS gp140 is gp120-gp41 bound by a disulfide bond NAbs 2G12, 2F5, IgG1b12, CD4 inducible 17b, and 19b bound to SOS gp140 better than uncleaved gp140 (gp140unc) and gp120 non-neutralizing MAbs 2.2B (binds to gp41 in gp140unc) and 23A (binds gp120) did not bind SOS gp140 2G12 complexes with SOS gp140 or with gp120 had a very unusual linear structure. Schulke *et al.* [2002] (antibody binding site definition and exposure, vaccine antigen design)
- 2G12: The antiviral response to intravenously administered MAbs 2F5 and 2G12 was evaluated in 7 HAART-naïve asymptomatic HIV-1 infected patients during a treatment period of 28 days. MAb therapy reduced plasma HIV RNA in 3/7 patients during the treatment period, and transiently reduced viral load in two more. CD4 counts were up in 3/7 through day 28, and transiently increased in three more. Vigorous complement activation was observed after 48/56 Ab infusions. Virus derived from 2/7 patients could be neutralized by 2G12, and escape from 2G12 was observed in both cases after infusion; one year after the infusion, isolates were again sensitive to 2G12. Stiegler *et al.* [2002] (complement, variant crossrecognition or cross-neutralization, escape, immunotherapy)
- 2G12: Passive immunization of neonate macaques with a combination of F105+2G12+2F5 conferred complete protection against oral challenge with SHIV-vpu+ or the combination b12+2G12+2F5 conferred partial protection against SHIV89.6 such combinations may be useful for prophylaxis at birth and against milk born transmission the synergistic combination of IgG1b12, 2G12, 2F5, and 4E10 neutralized a collection of HIV clade C primary isolates. Xu et al. [2002] (antibody interactions, immunoprophylaxis, mother-to-infant transmission)
- 2G12: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin stabilized oligomer gp140 delta683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and MAbs C11, A32, and 30D which did not bind the stabilized oligomer. Yang *et al.* [2002] (antibody binding site definition and exposure)
- 2G12: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002] (antibody binding site definition and exposure)

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- 2G12: SF162DeltaV2 is a virus that has a 30 amino acids deletion in the V2 loop that does not abrogate its infectivity but renders it highly susceptible to neutralization - when incorporated into a codon-optimized DNA vaccine with a CMV promoter and delivered by gene gun, SF162DeltaV2 gave higher neutralizing Ab titers against SF162 than did SF162 itself, and Abs that cross-neutralized non-homologous primary isolates were obtained only when SF162DeltaV2, but not intact SF162, was used as the immunogen - Control MAbs 2F5 and 2G12 could neutralize all of the following primary isolates: 91US056(R5), 92US714(R5), 92US660(R5), 92HT593(R5X4), and BZ167(R5X4), while after the first protein boost, the sera from two SF162DeltaV2 immunized macaques could neutralize 91US056(R5), 92US714(R5), 92US660(R5) and ADA(R5), but not 92HT593(R5X4) or 92US657(R5) - the pattern of cross-recognition shifted after the second boost. Barnett et al. [2001] (vaccine antigen design)
- 2G12: A combination of MAbs IgG1b12, 2F5, and 2G12 was given postnatally to four neonates macaques that were then challenged with highly pathogenic SHIV89.6P one of the four infants remained uninfected after oral challenge, two infants had no or a delayed CD4(+) T-cell decline. Hofmann-Lehmann et al. [2001] (immunoprophylaxis, mother-to-infant transmission)
- 2G12: Review of studies in macaques that have shown immune control of pathogenic SHIV viremia, improved clinical outcome, and protection, and the implications of the observations for HIV vaccines. Mascola & Nabel [2001] (review)
- 2G12: Moore and colleagues review structural aspects of gp120 and how they relate to antigenic domains, and review the data concerning the lack of a clear relationship between genetic subtype and serotype an exception exists for human MAb 2G12, which does not recognize CRF01 envelopes because of an unusual additional disulfide bond in the V4 loop region that appears to be unique to the subtype E, CRF01 gp120 protein. Moore *et al.* [2001] (antibody binding site definition and exposure, review)
- 2G12: Structural aspects of the interaction of neutralizing Abs with HIV-1 Env are reviewed Env essentially has three faces, one is largely inaccessible on the native trimer, and two that exposed but have low immunogenicity on primary viruses neutralization is suggested to occur by inhibition of the interaction between gp120 and the target cell membrane receptors as a result of steric hindrance and it is noted that the attachment of approximately 70 IgG molecules per virion is required for neutralization, which is equivalent to about one IgG molecule per spike the 2G12, 17b and b12 epitopes are discussed in detail although it is potently neutralizing, 2G12 does not interfere with CD4 and coreceptor binding, and this Ab specificity is uncommon in sera from HIV-1-infected individuals. Poignard *et al.* [2001] (antibody binding site definition and exposure, review)
- 2G12: Chloroquine reduces the HIV-1-infectivity of H9 IIIB cells, apparently through altering the conformation of envelope

   there is a reduction of reactivity of 2G12 to its epitope in chloroquine treated cultures. Savarino et al. [2001] (antibody binding site definition and exposure)

- 2G12: SHIV-HXBc2 is a neutralization sensitive non-pathogenic virus, and several *in vivo* passages through monkey's yielded highly pathogenic SHIV KU-1 HXBc2 and the KU-1 clone HXBc2P3.2 differ in 12 amino acids in gp160 substitutions in both gp120 and gp41 reduced the ability of sCD4, IgG1b12, F105 and AG1121 to Env achieve saturation and full occupancy, and neutralize KU-1 17b and 2F5 also bound less efficiently to HXBc2P3.2, although 2G12 was able to bind both comparably. Si *et al.* [2001]
- 2G12: A luciferase-reporter gene-expressing T-cell line was developed to facilitate neutralization and drug-sensitivity assays luciferase and p24 antigen neutralization titer end points were found comparable using NAb from sera from HIV+ donors, and MAbs 2F5, 2G12 and IgG1b12. Spenlehauer *et al.* [2001] (assay development)
- 2G12: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001] (antibody interactions)
- 2G12: Twenty HIV clade C isolates from five different countries were susceptible to neutralization by anti-clade B MAbs in a synergistic quadruple combination of mAbs IgG1b12, 2G12, 2F5, and 4E10. Xu et al. [2001] (antibody interactions, variant cross-recognition or cross-neutralization, subtype comparisons)
- 2G12: Neutralizing synergy between MAbs 1b12, 2G12 and 2F5 was studied using surface plasmon resonance to determine the binding kinetics for these three MAbs with respect to monomeric and oligomeric Env protein gp160 IIIB the 2G12 epitope is highly accessible on both monomeric and oligomeric Envs, 1b12 is highly accessible on monomers but not oligomers, and 2F5 on neither form binding of 2G12 exposes the 2F5 epitope on gp160 oligomers 2G12-gp160 oligomer interactions were best fitted to a two state model, with the first complex having a high association constant and fast dissociation, stabilized by conformational changes induced by the binding of a second MAb. Zeder-Lutz et al. [2001] (antibody binding site definition and exposure, antibody interactions, kinetics)
- 2G12: Neutralization synergy between anti-HIV NAbs b12, 2G12, 2F5, and 4E10 was studied a classic fixed-ratio method was used, as well as a method where one Ab was fixed at a low neutralization titer and the other was varied using primary isolates, a two-four fold enhancement of neutralization was observed with MAb pairs, and a ten-fold enhancement with a quadruple Ab combination no synergy was observed with any MAb pair in the neutralization of TCLA strain HXB2 there was no evidence for cooperativity of binding between b12 and 2G12 to envelope spikes expressed on the cell surface of TCLA or primary isolates. Zwick et al. [2001c] (antibody interactions)
- 2G12: A triple combination of 2F5, F105 and 2G12 effectively neutralized perinatal infection of macaque infants when challenged with SHIV-vpu+ the mean plasma half-life was 14.0

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- +/- 7.9 days, the longest of the three Abs. Baba *et al.* [2000] (immunoprophylaxis, mother-to-infant transmission)
- 2G12: To determine the antigenicity of virus killed by thermal and chemical inactivation, retention of conformation-dependent neutralization epitopes was examined, and exposure of CD4BS epitopes was found to be enhanced (MAbs IgG1b12, 205-46-9, and 205-43-1) binding to 2G12 and 447-52D epitopes was essentially unaltered the 17b CD4i epitope was also exposed. Grovit-Ferbas *et al.* [2000] (vaccine antigen design)
- 2G12: Because HIV-1 is most often transmitted across mucosal surfaces, the ability of passive transfer of infused HIVIG/2F5/2G12 to protect against mucosal exposure of macaques to pathogenic SHIV 89.6PD was studied HIVIG/2F5/2G12 protected 4/5 animals against vaginal challenge, 2F5/2G12 combined protected 2/5 animals, and 2G12 alone protected 2/4 animals in contrast, Mascola and coworkers had previously shown single MAbs could not protect against intervenous challenge Ab treated animals that got infected through vaginal innoculation had low viral loads and only modest declines in CD4 counts the infused Abs were detected in the nasal, vaginal, and oral mucosa. Mascola et al. [2000] (immunoprophylaxis, mucosal immunity)
- 2G12: Six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form 2G12 was an exception and could not neutralize MN in either form. Park et al. [2000]
- 2G12: A mini-review of observations of passive administration of IgG NAbs conferring protection against intervenous or vaginal SHIV challenge, that considers why IgG MAbs might protect against mucosal challenge. Robert-Guroff [2000] (immunoprophylaxis, mucosal immunity, review)
- 2G12: A Semliki Forest virus (SFV) expression system carrying BX08 Env was used to study the conformation of gp120 Env intracytoplasmic gp120 was recognized by the anti-V3 MAbs K24 and F5.5, while gp120 at the plasma membrane was detected only by conformation dependent MAbs 2G12, 670-D and 694/98D and not V3 MAbs expression in rat brain also showed that surface expressed Env was recognized only by the conformation-dependent Abs and not by anti-V3 Abs. Altmeyer et al. [1999]
- 2G12: rgp120 derived from a R5X4 subtype B virus was used to vaccinate healthy volunteers and the resulting sera were compared with sera from HIV-1 positive subjects and neutralizing MAbs 2G12 was able to bind with low affinity to the rgp120 monomer HIV-1 W61D. Beddows et al. [1999]
- 2G12: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 MAbs that

- bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley *et al.* [1999] (antibody binding site definition and exposure, vaccine antigen design)
- 2G12: Neutralization assays with rsCD4, MAbs, and serum samples from SHIV-infected macaques and HIV-1 infected individuals were used to characterize the antigenic properties of the env glycoprotein of six primary isolate-like or TCLA SHIV variants. 2G12 neutralized the five SHIV strains tested, HXBc2, KU2, 89.6, 89.6P and KB9, in MT-2 cells. Crawford *et al.* [1999] (variant cross-recognition or cross-neutralization)
- 2G12: Combinations of HIVIG, 2F5, 2G12 were administered in passive-transfer experiments 24 hours prior to challenge with pathogenic SHIV 89.6PD 3/6 animals given HIVIG/2F5/2G12 were completely protected, the others had reduced viremia and normal CD4 counts 1/3 monkeys given 2F5/2G12 showed transient infection, the other two had reduced viral load all monkeys that received HIVIG, 2F5, or 2G12 alone became infected and developed high-level plasma viremia, although animals that got HIVIG or 2G12 had a less profound CD4 T cell decline. Mascola *et al.* [1999] (antibody interactions)
- 2G12: A meeting summary presented results regarding neutralization –MAbs 2G12 and 2F5 tested for their ability to neutralize primary isolate infection of genetically engineered cell lines (cMAGI and others, presented by T. Matthews, A. Trkola, J. Bradac) an advantage of such cells lines over PBMCs is that markers (X-Gal) can be added for staining to simplify the assay the consensus of the meeting was that these engineered cell lines did not improve the sensitivity of detection of primary isolate neutralization D. Burton and J. Mascola presented results concerning passive immunization and protection of hu-PBL-SCID mice and macaques, respectively, and both found combinations of MAbs that were able to achieve 99% neutralization *in vitro* corresponded to efficacy *in vivo*. Montefiori & Evans [1999] (review)
- 2G12: Review of the neutralizing Ab response to HIV-1. Parren *et al.* [1999] (**review**)
- 2G12: Hu-PBL-SCID mice were infected with HIV-1s JRCSF and SF162 to study the effect of NAbs on an established infection no significant differences in the initial rate of decrease in viral load or the plateau levels of viral RNA between the b12 treated and control mice were seen in most of the Ab treated mice b12 escape mutants were observed with varying patterns of mutations a combination of b12, 2G12 and 2F5 protected 1/3 mice, and an isolate from one of the other two was resistant to neutralization by all three MAbs. Poignard *et al.* [1999] (antibody interactions, escape)
- 2G12: Post-exposure prophylaxis was effective when MAb 694/98-D was delivered 15 min post-exposure to HIV-1 LAI in hu-PBL-SCID mice, but declined to 50% if delivered 60 min post-exposure, and similar time constraints have been observed for HIVIG, 2F5 and 2G12, in contrast to MAb BAT123 that could protect when delivered 4 hours post infection. Andrus et al. [1998] (immunoprophylaxis)

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- 2G12: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein (Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer MAb 2G12 was the only exception to this, showing reduced binding efficiency. Binley et al. [1998] (antibody binding site definition and exposure)
- 2G12: Ab from gp120 vaccinated individuals prior to infection, who subsequently became HIV infected, could not achieve 90% neutralization of the primary virus by which the individuals were ultimately infected these viruses were not particularly refractive to neutralization, as determined by their susceptibility to neutralization by MAbs 2G12, IgG1b12, 2F5 and 447-52D. Connor *et al.* [1998]
- 2G12: Notes that 2G12 and 2F5, potent neutralizing antibodies, were identified by screening for cell surface (oligomeric Envelope) reactivity. Fouts *et al.* [1998] (antibody binding site definition and exposure)
- 2G12: Infection of dendritic cells cultured from CD14+ blood cells or from cadaveric human skin was blocked by neutralizing MAbs IgG1b12, or 2F5 and 2G12 delivered together, but not by control non-neutralizing anti-gp120 MAb 4.8D, indicating that NAbs could interrupt early mucosal transmission events. Frankel et al. [1998] (mucosal immunity)
- 2G12: The complete V, J and D(H) domain was sequenced unlike non-neutralizing anti-gp41 MAb 3D6, five neutralizing MAbs (2F5, 2G12, 1B1, 1F7, and 3D5) showed extensive somatic mutations giving evidence of persistent antigenic pressure over long periods 2G12 D(H) has the best homology to a D(H) segment between D3-22 and D4-23, a region not usually considered for heavy-chain rearrangement because it lacks associated recombination signals in the flanking regions, Kunert et al. suggest this may be why Abs that compete with 2G12 are rare. Kunert et al. [1998] (antibody sequence, variable domain)
- 2G12: Neutralization synergy was observed when the MAbs 694/98-D (V3), 2F5 (gp41), and 2G12 (gp120 discontinuous) were used in combination, and even greater neutralizing potential was seen with the addition of a fourth MAb, F105 (CD4 BS). Li *et al.* [1998] (antibody interactions)
- 2G12: Enhances Hx10 binding to CD4 positive or negative HeLa cells, but inhibited binding to CD4+ T-cell line A3.01
   neutralizes Hx10 infection of the HeLa cells. Mondor *et al.* [1998]
- 2G12: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren *et al.* [1998a] (antibody binding site definition and exposure)
- 2G12: MAbs 2G12, 2F5 and b12 are broadly neutralizing, as are some human polyconal sera, but this paper describes a set of primary isolates that are resistant to all three MAbs and 2 broadly neutralizing sera results indicate that resistance levels of pediatric isolates might be higher than adult isolates resistance in general did not seem to be conferred by a loss of binding affinity for gp120 or gp41, rather by a more global perturbation of oligomeric Envelope. Parren *et al.* [1998b] (variant cross-recognition or cross-neutralization)

- 2G12: In a study of the influence of the glycan at position 306 of the V3 loop on MAb recognition, 2G12 was found to neutralize an HIV-BRU mutant virus that lacks the V3 loop glycan and has a mutation at the tip of the loop more efficiently than it neutralizes HIV-BRU. Schonning *et al.* [1998] (antibody binding site definition and exposure)
  - 2G12: Does not compete with binding of MAb generated in response to gp120-CD4 complex, CG10. Sullivan *et al.* [1998b] (antibody interactions)
- 2G12: Induces complement-mediated lysis in MN but not primary isolates primary isolates are refractive to CML. Takefman et al. [1998] (complement, variant cross-recognition or cross-neutralization)
- 2G12: A wide range of neutralizing titers was observed that was independent of co-receptor usage. Trkola *et al.* [1998] (co-receptor, variant cross-recognition or cross-neutralization)
- 2G12: Summary of the implications of the crystal structure of gp120 combined with what is known about mutations that reduce NAb binding probable mechanism of neutralization by 2G12 is unknown, but dependent on proper glycosylation and 2G12 is predicted to be oriented toward the target cell when bound, so neutralization may be due to steric hindrance mutations in positions N 295, T 297, S 334, N 386, N 392 and N 397 HXBc2 (IIIB) decrease 2G12 binding, and the binding region is 25 angstroms from the CD4 binding site probably the Ab binds in part to carbohydrates, which may account for both its broad reactivity and the scarcity of Abs in the same competition group. Wyatt et al. [1998] (antibody binding site definition and exposure)
- 2G12: Review of the antigenic and receptor binding-domains of gp120 in relation to the structure of the molecule MAbs are discussed by category (anti-V2, anti-V3, CD4i, CD4BS...), however as 2G12 binds to a rarely immunogenic region, and it is dependent on glycosylation, it was discussed individually. Wyatt & Sodroski [1998] (review)
- 2G12: Review that discusses this MAb reacts with residues at the base of the V3 loop and V4, and most of the changes that reduce binding are glycosylation sites it is not clear whether the binding site is peptidic or direct carbohydrate. Burton & Montefiori [1997] (antibody binding site definition and exposure, review)
- 2G12: In a multilab evaluation of monoclonal antibodies, only IgG1b12, 2G12, and 2F5 could neutralize at least half of the 9 primary test isolates at a concentration of < 25 mug per ml for 90% viral inhibition neutralized 6 of 9 primary isolates. D'Souza *et al.* [1997] (variant cross-recognition or cross-neutralization)
- 2G12: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding 2G12 bound monomer, and weakly bound oligomer and neutralized JRFL. Fouts *et al.* [1997] (antibody binding site definition and exposure)
- 2G12: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB Env 2G12 was a strong neutralizer of SHIV-vpu+ all Ab combinations tested showed synergistic neutralization 2G12 has synergistic response with MAbs 694/98-D (anti-V3), 2F5, F105, and b12. Li *et al.* [1997] (antibody interactions)

**Env Antibodies HIV Antibodies Tables** 

- 2G12: Using concentrations of Abs achievable in vivo, the triple combination of 2F5, 2G12 and HIVIG was found to be synergistic to have the greatest breadth and magnitude of response against 15 clade B primary isolates. Mascola et al. [1997] (antibody interactions, variant cross-recognition or cross-neutralization)
- 2G12: A JRCSF variant that was selected for IgG1b12 resistance remained sensitive to MAbs 2G12 and 2F5, for combination therapy. Mo et al. [1997] (escape)
- 2G12: Review: MAbs 2F5, 2G12 and IgG1b12 have potential for use in combination with CD4-IgG2 as an immunotherapeutic or immunoprophylactic - homologous MAbs to these are rare in humans and vaccine strategies should consider including constructs that may enhance exposure of these MAbs' epitopes. Moore & Trkola [1997] (immunoprophylaxis, immunotherapv, review)
- 2G12: Neutralizes TCLA strains and primary isolates. Parren et al. [1997b] (variant cross-recognition or crossneutralization)
- 2G12: Viral binding inhibition by 2G12 was strongly correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5). Ugolini et al. [1997] (antibody binding site definition and exposure)
- 2G12: Neutralizes primary isolates, HXB2, and chimeric virus with gp120 from primary isolates in an HXB2 background. McKeating et al. [1996] (variant cross-recognition or crossneutralization)
- 2G12: Binding weakly enhanced by some anti-C1, -C4, -V3, and CD4 binding site MAbs - unusual in that 2G12 binding neither enhanced or inhibited the binding of other MAbs included in the study. Moore & Sodroski [1996] (antibody interactions)
- 2G12: Review: exceptional capacity to neutralize primary isolates in terms of both breadth and potency - one of three MAbs (IgG1b12, 2G12, and 2F5) generally accepted as having significant potency against primary isolates. Poignard et al. [1996b] (variant cross-recognition or cross-neutralization, review)
- 2G12: Review: Only four epitopes have been described which can stimulate a useful neutralizing response to a broad spectrum of primary isolates, represented by the binding sites of MAbs: 447-52-D, 2G12, Fab b12, and 2F5. Sattentau [1996] (review)
- 2G12: Conformationally sensitive epitope destroyed by mutations altering the N-linked glycosylation sites near the base of the V3 loop and the amino-terminal flank of the V4 loop. Trkola et al. [1996b] (antibody binding site definition and
- 2G12: Neutralizes JR-FL inhibits gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a] (**co-receptor**)
- 2G12: Review: binding site is distinct from CD4BS MAbs epitope and is unique among known gp120 MAbs, human or rodent. Moore & Ho [1995] (review)
- 2G12: Highly potent Cross-clade neutralizing activity. Trkola et al. [1995] (subtype comparisons)
- 2G12: Human MAb generated by electrofusion of PBL from HIV-1 + volunteers with CB-F7 cells. Buchacher et al. [1994] (antibody generation)

**No.** 800

MAb ID 30D **HXB2 Location** Env Author Location gp120

**Epitope** Neutralizing no

Immunogen

Species (Isotype)

References Yang et al. 2002

• 30D: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin—stabilized oligomer gp140Δ683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and C11, A32, and 30D which did not bind the stabilized oligomer. Yang et al. [2002]

No. 801

**MAb ID** 31710B

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human (IgG1)

References Alsmadi & Tilley 1998

• 31710B: A study of 6 anti-Env MAbs and their ability to bind or direct ADCC against target cells infected with IIIB, MN, SF-2, and RF – bound and directed lysis against all four strains. Alsmadi & Tilley [1998]

No. 802

MAb ID 38B5/C9

**HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 38B5/C9: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human  $IgG2\kappa$ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding these MAbs tended to be very cross-reactive but could not neutralize autologous SF162-38B5/C9 bound to most R5 and X4 B clade viruses, as well as one of two E clade viruses. He et al. [2002]

HIV Antibodies Tables Env Antibodies

No. 803

MAb ID 39H10/A11

**HXB2 Location** Env

Author Location gp120 (SF162)

Epitope
Subtype B
Neutralizing no
Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 39H10/A11: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2 $\kappa$  MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding—these MAbs tended to be very cross-reactive but could not neutralize autologous SF162—39H10/A11 bound to three R5 and three X4 B clade viruses, as well as two E clade viruses. He *et al.* [2002]

**No.** 804 **MAb ID** 3C9

**HXB2 Location** Env

Author Location gp120

Epitope
Subtype B
Neutralizing L

Immunogen vaccine

Strain: B clade SF2

Species (Isotype) mouse

References Kang et al. 1992

**Keywords** anti-idiotype, vaccine antigen design, variant cross-recognition or cross-neutralization

• C39: Murine antibodies were raised against human polyclonal antibodies against gp120, pooled from HIV-1 infected individuals. One anti-idiotype MAb was shown to bind to the CD4-binding site, and this MAb could raise anti-anti-idiotype antibodies when injected into cyngomolgous monkeys. The monkey MAbs neutralized laboratory strains MN, RF, and IIIB. Kang *et al.* [1992] (anti-idiotype, vaccine antigen design, variant cross-recognition or cross-neutralization)

No. 805

MAb ID 3D5

HXB2 Location Env

**Author Location** Env

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human

Research Contact Herman Katinger, Inst. Appl. Microbiol. University of Agricultural Science, Vienna, Aus-

tria

**References** Kunert *et al.* 1998; Purtscher *et al.* 1994; Buchacher *et al.* 1994

- 3D5: The complete V, J and D(H) domain was sequenced unlike non-neutralizing anti-gp41 MAb 3D6, five neutralizing MAbs (2F5, 2G12, 1B1, 1F7, and 3D5) showed extensive somatic mutations giving evidence of persistent antigenic pressure over long periods. Kunert et al. [1998]
- 3D5: Generated by electrofusion of PBL from HIV-1 positive volunteers with CB-F7 cells. Buchacher *et al.* [1994]

No. 806

MAb ID 3H6

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) mouse

References Pinter et al. 1995

- 3H6: There is another MAb with this ID that recognizes Rev.
- 3H6: Generated in response to virus grown in protein-free medium. Pinter et al. [1995]

**No.** 807

MAb ID 40D3/C11

**HXB2 Location** Env

**Author Location** gp120 (SF162)

**Epitope** 

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species** (**Isotype**) transgenic mouse ( $IgG2\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 40D3/C11: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2k MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding—these MAbs tended to be very cross-reactive but could not neutralize autologous SF162—40D3/C11 bound to most R5 and X4 B clade viruses, as well as one of two E clade viruses. He *et al.* [2002]

**No.** 808

MAb ID 49B11/A1

**HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** 

Subtype B

**Env Antibodies HIV Antibodies Tables** 

Neutralizing no Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad- Species (Isotype) transgenic mouse (IgG2κ)

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 49B11/A1: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human  $IgG2\kappa$ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding these MAbs tended to be very cross-reactive but could not neutralize autologous SF162—49B11/A1 bound to most R5 and X4 B clade viruses, as well as one of two E clade viruses. He et al. [2002]

No. 809

MAb ID 52G5/B9

**HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** Subtype B Neutralizing no Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad- Species (Isotype) transgenic mouse (IgG2κ) juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 52G5/B9: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human  $IgG2\kappa$ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120-11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding these MAbs tended to be very cross-reactive but could not neutralize autologous SF162-52G5/B9 bound to most R5 and X4 B clade viruses, as well as one of two E clade viruses. He et al. [2002]

**No.** 810 MAb ID 55E4/H1 **HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** Subtype B Neutralizing no Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 55E4/H1: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human  $IgG2\kappa$ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120-11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding these MAbs tended to be very cross-reactive but could not neutralize autologous SF162-55E4/H1 bound to most R5 and X4 B clade viruses, as well as one of two E clade viruses. He et al. [2002]

No. 811

MAb ID 56C4/C8

HXB2 Location Env

Author Location gp120 (SF162)

**Epitope** Subtype B Neutralizing no Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 56C4/C8: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human  $IgG2\kappa$ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120-11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their bindingthese MAbs tended to be very cross-reactive but could not neutralize autologous SF162-56C4/C8 bound to some R5 and X4 B clade viruses, as well as one of two E clade viruses. He et al. [2002]

No. 812

MAb ID 57B6/F1

**HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** Subtype B Neutralizing no Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI) **Species** (**Isotype**) transgenic mouse ( $IgG2\kappa$ )

HIV Antibodies Tables Env Antibodies

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org ing human genes allowing production of fully human IgG2 $\kappa$ 

References He et al. 2002

• 57B6/F1: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2k MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding—these MAbs tended to be very cross-reactive but could not neutralize autologous SF162—57B6/F1 bound some R5 and X4 B clade viruses, and no E clade viruses. He *et al.* [2002]

No. 813 MAb ID 57H5/D7

**HXB2 Location** Env

Author Location gp120 (SF162)

Epitope
Subtype B
Neutralizing no
Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 57H5/D7: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2k MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding—these MAbs tended to be very cross-reactive but could not neutralize autologous SF162—57H5/D7 bound to most R5 and X4 B clade viruses, as well as one of two E clade viruses. He et al. [2002]

**No.** 814

**MAb ID** 63G4/E2

**HXB2 Location** Env

**Author Location** gp120 (SF162)

Epitope Subtype B Neutralizing no Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species** (**Isotype**) transgenic mouse ( $IgG2\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research

Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 63G4/E2: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2κ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding—these MAbs tended to be very cross-reactive but could not neutralize autologous SF162—63G4/E2 bound to three R5 and three X4 B clade viruses, as well as two E clade viruses. He et al. [2002]

**No.** 815

MAb ID 65B12/C5

**HXB2 Location** Env

**Author Location** gp120 (SF162)

Epitope Subtype B Neutralizing no Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 65B12/C5: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2k MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—11 of the MAbs were conformation dependent, but did not block sCD4 binding—these MAbs were part of the same competition group, and enhanced binding of the CD4BS MAb 38G3/A9 and anti-CD4BS MAbs also enhanced their binding—these MAbs tended to be very cross-reactive but could not neutralize autologous SF162—65B12/C5 bound some R5 and X4 B clade viruses, and no E clade viruses. He *et al.* [2002]

**No.** 816

**MAb ID** 694/98D

**HXB2 Location** Env

**Author Location** Env (LAI)

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Ling et al. 2004

**Keywords** antibody binding site definition and exposure

• 694-98D: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of the V3 MAb 694-98D to its epitope was decreased by both thrombin and trypsin. Ling *et al.* [2004] (antibody binding site definition and exposure)

Env Antibodies HIV Antibodies Tables

**No.** 817

MAb ID 6D8

**HXB2 Location** Env

Author Location gp120 (21-85)

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype)

Research Contact Phil Berman

References Callahan et al. 1991

- Isolation of antibody.
- 6D8: Polyanionic polysaccharides were proposed to inhibit viral functions such as binding and syncytia formation through interations mediated through the local high positive charge density in the V3 loop. The binding of this N-term binding antibody is increased by dextransulfate, in contrast to anti-V3 antibodies that are inhibited. Callahan *et al.* [1991]

**No.** 818

MAb ID 6E10

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen

Species (Isotype)

Research Contact Phil Berman

References Callahan et al. 1991; Berman et al. 1991

- Isolation of antibody. Berman et al. [1991]
- 6E10: Polyanionic polysaccharides were proposed to inhibit viral functions such as binding and syncytia formation through interations mediated through the local high positive charge density in the V3 loop. The binding of this antibody is not inhibited by dextransulfate, in contrast to anti-V3 antibodies. Callahan *et al.* [1991]

**No.** 819

MAb ID 7-1054

**HXB2 Location** Env

Author Location gp36 (HIV-2)

**Epitope** 

Neutralizing no

**Immunogen** 

Species (Isotype) mouse

References Scheffel et al. 1999

 Binds HIV-2 gp36, used as a control in a study of group O MAbs. Scheffel et al. [1999]

**No.** 820

MAb ID 85G11/D8

**HXB2 Location** Env

Author Location gp120 (SF162)

Epitope

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: deglycosylated gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 85G11/D8: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2 $\kappa$  MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—three of the MAbs were conformation dependent, but did not block sCD4 binding and were part of the same competition group—these MAbs were all raised against a deglycosylated form of gp120—they could not neutralize autologous SF162 and bound some R5 and X4 B clade viruses, and no E clade viruses. He *et al.* [2002]

**No.** 821

**MAb ID** 87E4/A8

**HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** 

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: deglycosylated gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species** (**Isotype**) transgenic mouse ( $IgG2\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 87E4/A8: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2k MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—three of the MAbs were conformation dependent, but did not block sCD4 binding and were part of the same competition group—these MAbs were all raised against a deglycosylated form of gp120—they could not neutralize autologous SF162 and bound some R5 and X4 B clade viruses, and no E clade viruses. He *et al.* [2002]

**No.** 822

**MAb ID** 97B1/E8

**HXB2 Location** Env

**Author Location** gp120 (SF162)

**Epitope** 

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: deglycosylated gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

97B1/E8: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2κ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—three of the MAbs were conformation dependent, but

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did not block sCD4 binding and were part of the same compe- Author Location gp41 (717–751) tition group—these MAbs were all raised against a deglycosylated form of gp120—they could not neutralize autologous SF162 and bound some R5 and X4 B clade viruses, and no E clade viruses. He et al. [2002]

No. 823

MAb ID A9

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: chimeric GM-CSF Strain: B clade IIIB HIV component: gp120 Adjuvant: GM-CSF

Species (Isotype) mouse (IgG1)

References del Real et al. 1999

• A9: Murine antibody response to the chimeric construction granulocyte-macrophage colony stimulating factor GM-CSF/gp120 was tested, MAbs from normal mice were gp120 specific, MAbs from nude mice bound gp120 but were polyreactive, and from reconstituted mice were half way between - the anti-gp120 response used a high frequency of VH81X, VHQ52, and VH7183 genes, a family used during fetal life and associated with autoimmunity – A9 was a gp120 from a BALBc reconstructed nude mouse and had VH gene 7183-2. del Real et al. [1999]

**No.** 824

MAb ID ADP421 polyclonal

**HXB2 Location** Env

Author Location Env

**Epitope** 

Subtype A

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) rabbit

References Jeffs et al. 2004

**Keywords** subtype comparisons, vaccine antigen design • ADP421: A set of oligomeric envelope proteins were made from six primary isolates for potenial use as vaccine antigens: 92/UG/037 (clade A), HAN2/2 (clade B), 92/BR25/025 (clade C), 92/UG/021 (clade D), 93/BR/029 (clade F) and MVP5180 (clade O). This was one of a panel of MAbs used to explore folding and exposure of well characterized epitopes. The clade C isolate BR25 is apparently misfolded, as conformationdependent antibodies did not bind to it. ADP421 is a polyclonal rabbit sera raised against CHO-derived IIIB gp120. ADP421 bound to antigens from all clades A-F, as well as group O. Polyclonal sera raised in rabbits against these antigens crossbound the other antigens, but none of the sera had neutralizing activity. Jeffs et al. [2004] (vaccine antigen design, subtype comparisons)

No. 825 MAb ID AG10H9 **HXB2 Location** Env

**Epitope** 

Neutralizing

**Immunogen** 

Species (Isotype)

Research Contact BabCO

References Ohagen et al. 2003

Keywords brain/CSF, variant cross-recognition or crossneutralization

• AG10H9: Env genes derived from uncultured brain biopsy samples from four HIV-1 infected patients with late-stage AIDS were compared to env genes from PBMC samples. Brain isolates did not differ in the total number or positions of Nglycosylation sites, patterns of coreceptor usage, or ability to be recognized by gp160 and gp41 MAbs. AG10H9 recognized most variants gp41 and gp160 from 3/4 individuals by WB, but not the 4th. Ohagen et al. [2003] (brain/CSF, variant cross-recognition or cross-neutralization)

No. 826

MAb ID AH48

**HXB2 Location** Env

Author Location gp120 (V3)

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Zwick et al. 2003

Keywords antibody generation, antibody interactions

• AH-48: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. AH48 is a novel anti-V3 Fab first used in this study. Zwick et al. [2003] (antibody generation, antibody interactions)

No. 827

MAb ID B4

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: chimeric GM-CSF Strain: B

clade IIIB HIV component: gp120

Species (Isotype) mouse (IgM)

References del Real et al. 1999

• B4: Murine antibody response to the chimeric construction granulocyte-macrophage colony stimulating factor GM-CSF/gp120 was tested, MAbs from normal mice were gp120 specific, MAbs from nude mice bound gp120 but were polyreactive, and from reconstituted mice were half way between - the anti-gp120 response used a high frequency of VH81X, Env Antibodies HIV Antibodies Tables

VHQ52, and VH7183 genes, a family used during fetal life and associated with autoimmunity – B4 was an anti-gp120 from a BALBc reconstructed nude mouse and had VH gene J606. del Real *et al.* [1999]

No. 828

MAb ID B5

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope Neutralizing** 

T.....

Immunogen vaccine

Vector/Type: chimeric GM-CSF Strain: B clade IIIB HIV component: gp120 Adju-

vant: GM-CSF

Species (Isotype) mouse (IgG1)

References del Real et al. 1999

B5: Murine antibody response to the chimeric construction granulocyte-macrophage colony stimulating factor GM-CSF/gp120 was tested, MAbs from normal mice were gp120 specific, MAbs from nude mice bound gp120 but were polyreactive, and from reconstituted mice were half way between – the anti-gp120 response used a high frequency of VH81X, VHQ52, and VH7183 genes, a family used during fetal life and associated with autoimmunity – B5 was a gp120 specific MAb from a BALBc mouse and had VH gene J558. del Real et al. [1999]

**No.** 829

MAb ID B6

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope Neutralizing** 

Immunogen vaccine

*Vector/Type:* chimeric GM-CSF *Strain:* B clade IIIB *HIV component:* gp120

Species (Isotype) mouse (IgM)

References del Real et al. 1999

• B6: Murine antibody response to the chimeric construction granulocyte-macrophage colony stimulating factor GM-CSF/gp120 was tested, MAbs from normal mice were gp120 specific, MAbs from nude mice bound gp120 but were polyreactive, and from reconstituted mice were half way between – the anti-gp120 response used a high frequency of VH81X, VHQ52, and VH7183 genes, a family used during fetal life and associated with autoimmunity – B6 was a gp120 from a BALBc reconstructed nude mouse and had VH gene J558. del Real *et al.* [1999]

**No.** 830

MAb ID BAT267

**HXB2 Location** Env

Author Location gp120

Epitope

Neutralizing L

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Fung et al. 1987

No. 831

MAb ID BAT401

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B

clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Fung et al. 1987

No. 832

MAb ID BAT509

HXB2 Location Env

**Author Location** gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: inactivated HIV Strain: B

clade IIIB HIV component: HIV-1

Species (Isotype) mouse (IgG1)

References Fung et al. 1987

**No.** 833

MAb ID C31

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

References Boyer et al. 1991

C31: Broadly-reactive group specific MAb – high yield cultivation of human MAb. Boyer et al. [1991]

**No.** 834

MAb ID CO11

**HXB2 Location** Env

Author Location gp120 (V3)

Epitope

Neutralizing

Immunogen

Species (Isotype)

Ab Type gp120 V3

Research Contact James Robinson, Tulane University, New Or-

leans, LA, USA

References Haynes et al. 2005; Pantophlet et al. 2004;

Grundner et al. 2002

Keywords antibody binding site definition and exposure,

vaccine antigen design

 CO11: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. CO11 has no indication of polyspecific **HIV Antibodies Tables Env Antibodies** 

autoreactivity. Haynes et al. [2005] (antibody binding site • D12: MAbs D10 and D12 are very easily blocked by human definition and exposure)

• CO11: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only 3 additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including CO11. Pantophlet et al. [2004] (vaccine antigen design)

No. 835

MAb ID D1

**HXB2 Location** Env

Author Location gp41 (IIIB)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

References Otteken et al. 1996

• D1: MAbs D1, D16, had T37 bind to oligomeric gp160 equally well – pulse label experiments of MAb binding to noncleavable gp160 revealed that these MAbs bound with a delay, epitopes forming with a half life of 30 min. Otteken et al. [1996]

No. 836

MAb ID D12

**HXB2 Location** Env

Author Location gp41 (IIIB)

**Epitope** Neutralizing L Immunogen vaccine

> Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Research Contact Patricia Earl and Christopher Broder, NIH

References Yang et al. 2000; LaBranche et al. 1999; Otteken et al. 1996; Earl et al. 1997; Richardson

et al. 1996; Broder et al. 1994; Earl et al. 1994

**Keywords** antibody binding site definition and exposure, antibody generation, antibody interactions, vaccine antigen design

- D12: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) – gp41 MAbs T4, D12, T3, and D50 bound less efficiently to gp140-GNC4 than did pooled sera, but T4 and D12 recognized the gp140-GNC4 timer equivalently to gp140(-), and T3 and D50 recognized the trimer at greater levels than gp140(-). Yang et al. [2000] (vaccine antigen design)
- D12: D12 was used in WB of HIV-1 transmembrane proteins in a study which showed that determinants of HIV-1 CD4 independence map outside regions required for coreceptor specificity – IIIBx, a CD4-independent variant of IIIB, has a truncated gp41. LaBranche et al. [1999]

sera from HIV+ individuals. Earl et al. [1997]

- D12: MAbs D4, D10, D11, D12, and D41 all bind only to complete oligomer - pulse label experiments of MAb binding to noncleavable gp160 revealed that these MAbs bound with a delay, epitopes forming with a half life of 30 min. Otteken et al. [1996] (antibody binding site definition and exposure)
- D12: This antibody was blocked more strongly by human sera than other anti-gp41 MAbs (D20, D43, D61, and T4) in a oligomeric ELISA assay. Richardson et al. [1996] (antibody interactions)
- D12: One of 18 MAbs (e. g. D4 and D40) that bind to a conformation-dependent epitope in gp41 that bind preferentially, but not exclusively, to oligomers – neutralizes IIIB and SF2. Broder et al. [1994] (antibody binding site definition and exposure)
- D12: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994] (antibody generation)

No. 837

MAb ID D16

**HXB2 Location** Env

**Author Location** gp41 (IIIB)

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein HIV component:

dimeric Env

Species (Isotype) mouse (IgG)

Research Contact Patricia Earl and Christopher Broder, NIH

References Earl et al. 1997; Weissenhorn et al. 1996; Earl et al. 1994

- D16: One of eleven MAbs (D16, D17, D31, D36, D37, D40, D44, D55, D59, T37, and T45) that are conformation dependent and that can block the binding of the MAb D50 that binds to the linear peptide gp41(642-665) – reactive with 9/10 HIV-1 strains all except HIV-1 ADA, which has the change E659D and E662A that may result in the loss of binding (ELLE to DLLA). Earl *et al*. [1997]
- D16: Precipitates both oligomeric gp140 and soluble monomeric gp41(21-166)that lacks the fusion peptide and membrane anchor, along with MAbs D16, D38, D40, D41, and D54. Weissenhorn et al. [1996]
- D16: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

No. 838

MAb ID D4

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: chimeric GM-CSF Strain: B

clade IIIB HIV component: gp120

Species (Isotype) mouse (IgG1)

References del Real et al. 1999

• D4: Murine antibody response to the chimeric construction granulocyte-macrophage colony stimulating factor GM-CSF/gp120 was tested, MAbs from normal mice were gp120 specific, MAbs from nude mice bound gp120 but were polyreactive, and from reconstituted mice were half way between - the anti-gp120 response used a high frequency of VH81X, VHQ52, and VH7183 genes, a family used during fetal life and associated with autoimmunity – D4 was a gp120 from a BALBc reconstructed nude mouse and had VH gene J558. del Real et al. [1999]

**No.** 839 MAb ID D43 **HXB2 Location** Env

Author Location gp41 (HXB2)

**Epitope** Subtype B Neutralizing Immunogen vaccine

> protein HIV component: Species (Isotype) human Vector/Type:

dimeric Env

Species (Isotype) mouse (IgG)

Research Contact Patricia Earl and Christopher Broder, NIH References Earl et al. 1997; Richardson et al. 1996; Earl et al. 1994

- D43: Partially conformation dependent doesn't bind to short peptides, but does bind to the region spanning 641-683 - binding can be blocked by MAbs T3, D38 and D45 - MAbs in this competition group reacted with 9/10 HIV-1 strains, not binding to JRFL. Earl et al. [1997]
- D43: This is a linear gp41 epitope, mapping in the region 635-678 - human sera blocked binding in oligomeric ELISA assay to a similar extent for gp41 MAbs D20, D43, D61, and T4. Richardson et al. [1996]
- D43: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 840

MAb ID F223

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG3 $\lambda$ )

References Cavacini et al. 1999

• F223: binds to HIV-1 gp120 and to uninfected lymphocytes binding to a 159-kd auto-antigen expressed on most B cells and a small fraction of T and NK cells - the antibody enhances HIV-1 infection in a complement-dependent manner -F223 light chains have a strong homology with VLgamma2, the heavy chain to the germline gene VH3-H.11 - N-linked carbohydrates are key for recognition of both gp120 and the autoantigen – MAb 3D6 also uses VH3 and has autoreactivity. Cavacini *et al*. [1999]

No. 841 MAb ID F285 **HXB2 Location** Env

**Author Location** Env

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1)

References Wisnewski et al. 1996; Wisnewski et al. 1995 • F285: F285 is V H1 – V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996]

No. 842

MAb ID F424

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

References Ferrantelli et al. 2004a

Keywords variant cross-recognition crossneutralization

• F424: Nabs against HIV-1 M group isolates were tested for their ability to neutralize 6 randomly selected HIV-1 O group strains. F424 is noted to be a MAb derived from a subtype B infected individual, that binds to an undefined epitope in gp120 and can neutralize some M group viruses, but it was not particularly effective at neutralization of the O group viruses tested. Ferrantelli et al. [2004a] (variant cross-recognition or cross-neutralization)

No. 843

MAb ID F425 B4e8 (F425-B4e8)

**HXB2 Location** Env

**Author Location** gp120 (V3)

**Epitope** 

**Neutralizing** 

**Immunogen** 

Species (Isotype)

Ab Type gp120 V3

Research Contact L. Cavacini

**References** Selvarajah et al. 2005; Pantophlet et al. 2004; Cavacini et al. 2003

Keywords vaccine antigen design, vaccine-specific epi-

tope characteristics

• F425 B4e8: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. V3 MAbs (447-52D, 19b, F245-B4e8 and 39F) bound to the GDMR antigen, but either did not bind or had diminished binding to mCHO. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)

• F425 B4e8: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including F425 B4e8. This MAb bound to the initial construct, but introduction of glycosylation sites at positions 320 and 325 inhibited binding. Pantophlet et al. [2004]

No. 844 MAb ID F7 HXB2 Location Env Author Location gp120 (IIIB) **Epitope** Neutralizing Immunogen vaccine

> Vector/Type: chimeric GM-CSF Strain: B clade IIIB HIV component: gp120 Adju-

vant: GM-CSF

Species (Isotype) mouse (IgG1)

References del Real et al. 1999

• F7: Murine antibody response to the chimeric construction granulocyte-macrophage colony stimulating factor GM-CSF/gp120 was tested, MAbs from normal mice were gp120 specific, MAbs from nude mice bound gp120 but were polyreactive, and from reconstituted mice were half way between - the anti-gp120 response used a high frequency of VH81X, VHQ52, and VH7183 genes, a family used during fetal life and associated with autoimmunity – F7 was a gp120 specific MAb from a BALBc mouse and had VH gene 7183(81X), previously found expressed only in fetal liver. del Real et al. [1999]

No. 845 MAb ID Fab A12 **HXB2 Location** Env Author Location gp41 (LAI) **Epitope** Subtype B Neutralizing no Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ )

References Binley et al. 1996 • Fab A12: Uncharacterized epitope - variable regions sequenced. Binley et al. [1996]

**No.** 846 MAb ID Fab A2 **HXB2 Location** Env Author Location gp41 (LAI) **Epitope** Subtype B Neutralizing no

Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\lambda$ ) References Binley et al. 1996 • Fab A2: Uncharacterized epitope – variable regions sequenced. Binley et al. [1996]

No. 847 MAb ID Fab L9 **HXB2 Location** Env Author Location gp41 (LAI) **Epitope** Subtype B Neutralizing no Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ )

References Binley et al. 1996

• Fab L9: Uncharacterized epitope – variable regions sequenced. Binley et al. [1996]

No. 848 MAb ID G12 **HXB2 Location** Env Author Location gp120 (IIIB) **Epitope Neutralizing** Immunogen vaccine

> Vector/Type: chimeric GM-CSF Strain: B clade IIIB HIV component: gp120

Species (Isotype) mouse (IgM)

References del Real et al. 1999

· G12: Murine antibody response to the chimeric construction granulocyte-macrophage colony stimulating factor GM-CSF/gp120 was tested, MAbs from normal mice were gp120 specific, MAbs from nude mice bound gp120 but were polyreactive, and from reconstituted mice were half way between - the anti-gp120 response used a high frequency of VH81X, VHQ52, and VH7183 genes, a family used during fetal life and associated with autoimmunity - G12 was a gp120 from a BALBc reconstructed nude mouse and had VH gene 7183-6. del Real et al. [1999]

No. 849 MAb ID G2 **HXB2 Location** Env Author Location gp120 (IIIB) **Epitope Neutralizing** Immunogen vaccine

> Vector/Type: chimeric GM-CSF Strain: B clade IIIB HIV component: gp120

Species (Isotype) mouse (IgM)

References del Real et al. 1999

• G2: Murine antibody response to the chimeric construction granulocyte-macrophage colony stimulating factor GM-CSF/gp120 was tested, MAbs from normal mice were gp120 specific, MAbs from nude mice bound gp120 but were polyreactive, and from reconstituted mice were half way between - the anti-gp120 response used a high frequency of VH81X, VHQ52, and VH7183 genes, a family used during fetal life and associated with autoimmunity - G2 was a gp120 from a BALBc reconstructed nude mouse and had VH gene Q52. del Real et al. [1999]

**No.** 850

MAb ID H2

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

**Neutralizing** 

**Immunogen** 

**Species** (**Isotype**) human ( $IgM\kappa$ )

Research Contact BioInvent, Lund, Sweden, commercial

References Muller et al. 1991

• H2: Anti-idiotypic MAbs (10B3 and 2All) against MAb H2 were generated by immunization of BALBc mice with H2 they also react with seropositive sera. Muller et al. [1991]

No. 851

MAb ID H8

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope Neutralizing** 

Immunogen vaccine

clade IIIB HIV component: gp120

**Species (Isotype)** mouse (IgM)

References del Real et al. 1999

• H8: Murine antibody response to the chimeric construction granulocyte-macrophage colony stimulating factor GM-CSF/gp120 was tested, MAbs from normal mice were gp120 specific, MAbs from nude mice bound gp120 but were polyreactive, and from reconstituted mice were half way between - the anti-gp120 response used a high frequency of VH81X, VHQ52, and VH7183 genes, a family used during fetal life and associated with autoimmunity - H8 was a gp120 from a BALBc reconstructed nude mouse and had VH gene Q52. del Real et al. [1999]

No. 852

MAb ID HBW4

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

References Wisnewski et al. 1996; Wisnewski et al. 1995; Moran et al. 1993

- HBW4: HBW4 is V H2 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996]
- HBW4: Heavy (V HII) and light (V lambdaII) chain sequenced. Moran et al. [1993]

**No.** 853

MAb ID HIVIG

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** P

Immunogen HIV-1 infection

Species (Isotype) human

References Nichols et al. 2002

• NYBC-HIVIG derived from patients with high NAb titers and NABI-HIVIG derived from patients with high anti-p24 Ab titers were compared in neutralizing assay against a panel of six primary isolates - both could neutralize all isolates tested but the NYBC-HIVIG dose required for 50% neutralization was of 3.2 fold lower, showing source plasmas influence the effective concentration of NAb present in HIVIG. Nichols et al. [2002]

No. 854

MAb ID IVI-4G6

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing

Immunogen vaccine

Species (Isotype) mouse (IgG2b)

Research Contact K. Miyakoshi (Feji-Rebio Co, Tokyo, Japan)

References Yin et al. 2001

Vector/Type: chimeric GM-CSF Strain: B • IVI-4G6: A bi-specific Ab (BFA) was made by combining Fab fragments of gp41-specific MAb IVI-4G6 and CD3-specific Mab UCHT1 - the BFA suppressed HIV-1 propagation culture and eliminated latently infected cells. Yin et al. [2001]

No. 855

MAb ID IgA6/30λ

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing ves

Immunogen HIV-1 exposed seronegative

Species (Isotype) human

References Berry et al. 2003

Country Kenya

**Keywords** antibody generation, antibody sequence, variable domain, genital and mucosal immu-

nity, HIV exposed persistently seronegative

(HEPS)

• A panel of anti-gp120 single-chain variable fragment (scFv) Ab was isolated from cervical B lymphocytes of unexposed uninfected Kenyan prostitutes. These Abs recognize gp120 in ELISA and using flow cytometry. IgG1b12 does not inhibit binding of the new clones to HIV, so the epitopes are distinct. Sequencing of the V genes of the scFv clones show they are unique. Berry et al. [2003] (antibody generation, genital and mucosal immunity, HIV exposed persistently seronegative (HEPS), antibody sequence, variable domain)

No. 856

MAb ID IgA6/5k

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing yes

Immunogen HIV-1 exposed seronegative

Species (Isotype) human

References Berry et al. 2003

Country Kenya

**Keywords** antibody generation, antibody sequence, variable domain, genital and mucosal immunity, HIV exposed persistently seronegative (HEPS)

A panel of anti-gp120 single-chain variable fragment (scFv)
 Ab was isolated from cervical B lymphocytes of unexposed
 uninfected Kenyan prostitutes. These Abs recognize gp120 in
 ELISA and using flow cytometry. IgG1b12 does not inhibit
 binding of the new clones to HIV, so the epitopes are distinct.
 Sequencing of the V genes of the scFv clones show they are
 unique. Berry et al. [2003] (antibody generation, genital and
 mucosal immunity, HIV exposed persistently seronegative
 (HEPS), antibody sequence, variable domain)

No. 857

MAb ID IgA6/L4

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing yes

Immunogen HIV-1 exposed seronegative

Species (Isotype) human

References Berry et al. 2003

Country Kenya

**Keywords** antibody generation, antibody sequence, variable domain, genital and mucosal immunity, HIV exposed persistently seronegative (HEPS)

• A panel of anti-gp120 single-chain variable fragment (scFv) Ab was isolated from cervical B lymphocytes of unexposed uninfected Kenyan prostitutes. These Abs recognize gp120 in ELISA and using flow cytometry. IgG1b12 does not inhibit binding of the new clones to HIV, so the epitopes are distinct. IgA6/4L is neutralizing. Sequencing of the V genes of the scFv clones show they are unique. Berry et al. [2003] (antibody generation, genital and mucosal immunity, HIV exposed persistently seronegative (HEPS), antibody sequence, variable domain)

No. 858

MAb ID K14

HXB2 Location Env

**Author Location** gp41

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype) human (IgG1)

**References** Schutten *et al.* 1997; Schutten *et al.* 1996; Schutten *et al.* 1995b; Schutten *et al.* 1995a; Teeuwsen *et al.* 1990

- K14: In a study of NSI and SI virus neutralization, K14 did not influence viral entry. Schutten *et al.* [1997]
- K14: Reduced affinity for both SI and NSI viruses relative to MAb MN215, failed to neutralize SI strain. Schutten *et al.* [1995b]
- K14: Did not bind to peptides spanning gp41, but it does not react with Env deletion mutant 643-692 does not react with HIV-2– competition experiments showed this was an immunodominant conserved epitope in HIV-1 positive sera from Europe and Africa. Teeuwsen *et al.* [1990]

**No.** 859

MAb ID KU32

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing

**Immunogen** 

Species (Isotype) human

References Haynes et al. 2005

Keywords antibody binding site definition and exposure • KU32: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. KU32 was noted to have some polyspecific autoreactivity in the text, but it was not clear how this was manifested from the results. Haynes *et al.* [2005] (antibody

binding site definition and exposure)

No. 860

MAb ID M25

**HXB2 Location** Env

Author Location gp41

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: purified HIV-1

**Species (Isotype)** mouse ( $IgG\kappa$ )

**References** Watkins *et al.* 1996; di Marzo Veronese *et al.* 

 M25: heavy and light chains cloned and sequenced – binding requires heavy and light chain in combination, in contrast to M77. Watkins et al. [1996]

No. 861

MAb ID MAG 6B

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Research Contact C. Y. Kang, IDEC Inc

References Kang et al. 1994

 MAG 6B: Amino acid substitutions that reduce binding 10 fold: 256 S/Y, 257 T/R or G or A, 262 N/T, 368 D/R or T, 370 E/R or Q, 381 E/P, 384 Y/E, 421 K/L, 475 M/S, 477 D/V. Kang et al. [1994]

**No.** 862

MAb ID MO28

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

Neutralizing no

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgM)

References Ohlin et al. 1989

• MO28: This antibody was raised by *in vitro* stimulation with a recombinant Env penv9 – the discontinuous epitope involves hydrophobic regions 632-646, 677-681 and 687-691, proximal to and spanning the transmembrane region – this specificity is unusual in HIV-1 positive sera. Ohlin *et al.* [1989]

No. 863

MAb ID MO30

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing no

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgM)

References Ohlin et al. 1989

• MO30: This antibody was raised by *in vitro* stimulation with a recombinant Env penv9 – the discontinuous epitope involves hydrophobic regions 632-646, 677-681 and 687-691, proximal to and spanning the transmembrane region – this specificity is unusual in HIV-1 positive sera. Ohlin *et al.* [1989]

**No.** 864

MAb ID MO43

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

Neutralizing no

Immunogen in vitro stimulation or selection

Species (Isotype) human (IgM)

References Ohlin et al. 1989

• MO43: This antibody was raised by *in vitro* stimulation with a recombinant Env penv9 – the discontinuous epitope of MO43 involves hydrophobic regions 632-646, 677-681 and 687-691, proximal to and spanning the transmembrane region – this specificity is unusual in HIV-1 positive sera. Ohlin *et al.* [1989]

No. 865

MAb ID N2-4

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG1 $\kappa$ )

Research Contact Evan Hersh and Yoh-Ichi Matsumoto

References Robinson et al. 1990b

- N2-4: NIH AIDS Research and Reference Reagent Program: 528
- N2-4: No enhancing activity for HIV-1 IIIB. Robinson *et al.* [1990b]

**No.** 866

MAb ID N70-2.3a

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Research Contact James Robinson, Tulane University, LA

References Takeda et al. 1992; Robinson et al. 1990a

- N70-2.3a: Fc receptor mediated enhancement of HIV-1 infection binds a conformational site in the carboxyl half of gp120, distinct from 1.5e. Takeda *et al.* [1992]
- N70-2.3a: Broad reactivity. Robinson et al. [1990a]

**No.** 867

**MAb ID** P43110

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen

Species (Isotype)

Research Contact Advanced Biosciences (Kensington, MD)

**References** VanCott *et al.* 1995; di Marzo Veronese *et al.* 1992

P43110: Does not recognized denatured form of the gp120 protein. VanCott et al. [1995]

No. 868

MAb ID P5-3

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Research Contact Evan Hersh and Yoh-Ichi Matsumoto

References Pincus et al. 1991; Robinson et al. 1990b

- P5-3: NIH AIDS Research and Reference Reagent Program: 378
- P5-3: Poor immunotoxin activity when coupled to RAC isotype specified as: IgG3lambda. Pincus *et al.* [1991]
- P5-3: No enhancing activity for HIV-1 IIIB. Robinson *et al.* [1990b]

No. 869

MAb ID T15G1

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype)

References Binley et al. 1999

• T15G1: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits – a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen – SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 – SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 – nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly

induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999]

**No.** 870

MAb ID T20

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

> References Sugiura et al. 1999; Otteken et al. 1996; Earl et al. 1994

- T20: A comparison of 25 gp120 specific, conformation dependent MAbs was done - T20 is part of a group of MAbs labeled AII - all AII MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, and could only partially blocked CD4 binding. Sugiura et al. [1999]
- T20: Pulse label experiments of 4 MAbs (D20, D27, T20, and T22) binding to noncleavable gp140 revealed that these anti-CD4BS MAbs bound with a delay, and that the epitope formed with a t 1/2 of about 10 minutes. Otteken et al. [1996]
- T20: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 871

MAb ID T27

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- T27: A comparison of 25 gp120 specific, conformation dependent MAbs was done - T27 is part of a group of MAbs labeled AII - all AII MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, and could only partially blocked CD4 binding. Sugiura et al. [1999]
- T27: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

No. 872

MAb ID T3

**HXB2 Location** Env

Author Location gp41 (HXB2)

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: tetrameric Env HIV compo-

nent: Env

Species (Isotype) mouse (IgG)

References Yang et al. 2000; Zwick et al. 2001b; Earl et al. 1997; Earl et al. 1994

- T3: T3 partially competes with MAb Z13, but not MAb 4E10, both of which bind to gp41 proximally to the 2F5 epitope and have a broad neutralizing potential. Zwick et al. [2001b]
- T3: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) – gp41 MAbs T4, D12, T3, and D50 bound less efficiently to gp140-GNC4 than did pooled sera, but T4 and D12 recognized the gp140-GNC4 timer equivalently to gp140(-), and T3 and D50 recognized the trimer at greater levels than gp140(-). Yang et al. [2000]
- T3: Partially conformation dependent doesn't bind to short peptides, but does bind to the region spanning 641-683 - binding can be blocked by MAbs D43, D38 and D45 - MAbs in this competition group reacted with 9/10 HIV-1 strains, not binding to JRFL. Earl et al. [1997]
- T3: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

No. 873

MAb ID T30

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: tetrameric Env HIV compo-

nent: Env

Species (Isotype) mouse

Research Contact C. Broder

References Ohagen et al. 2003; Earl et al. 1997; Earl et al.

Keywords antibody binding site definition and exposure, antibody generation, brain/CSF, escape

- T30: Env genes derived from uncultured brain biopsy samples from four HIV-1 infected patients with late-stage AIDS were compared to env genes from PBMC samples. Brain isolates did not differ in the total number or positions of N-glycosylation sites, patterns of coreceptor usage, or ability to be recognized by gp160 and gp41 MAbs. T30 recognized most variants (10/13) gp41 by WB, and all of the gp160s. Ohagen et al. [2003] (brain/CSF, escape)
- T30: Binds in the region 580 to 640, but does not bind to peptides spanning this region – binding depends on N-linked glycosylation of Asn 616 – no other antibody tested inhibited binding, but binding could be inhibited by sera from HIV+ individuals. Earl et al. [1997] (antibody binding site definition and exposure)
- T30: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994] (antibody generation)

No. 874

MAb ID T4

**HXB2 Location** Env

Author Location gp41 (IIIB)

**Epitope** Neutralizing L Immunogen vaccine

> Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

References Srivastava et al. 2002; Yang et al. 2000; Stamatatos et al. 2000; Binley et al. 1999; Earl et al. 1997; Otteken et al. 1996; Weissenhorn et al. 1996; Richardson et al. 1996; Broder et al. 1994; Earl et al. 1994

Keywords antibody binding site definition and exposure, antibody generation, antibody interactions, vaccine antigen design

- T4: Oligomeric gp140 (o-gp140) derived from R5 primary isolate US4 was characterized for use as a vaccine reagent antigen capture ELISA was used to compare the antigenicity of gp120 and o-gp140 using a panel of well characterized MAbs - T4 recognized o-gp140. Srivastava et al. [2002] (antibody binding site definition and exposure)
- T4: Soluble gp140 derived from SF162, a neutralizationresistant primary isolate, and SF162AV2 a neutralizationsusceptible isolate with 30 amino acids deleted from the V2 loop, were generated with or without the gp120-gp41 cleavage site intact – all forms are recognized by oligomer-specific MAb T4 and show enhanced binding of CD4i MAb 17b when sCD4 is bound - the fused forms are less efficiently recognized than the cleaved forms by polyclonal neutralizing sera from HIVinfected patients - the V3 loop is more exposed on the fused form. Stamatatos et al. [2000] (vaccine antigen design)
- T4: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) - gp41 MAbs T4, D12, T3, and D50 bound less efficiently to gp140-GNC4 than did pooled sera, but T4 and D12 recognized the gp140-GNC4 timer equivalently to gp140(-), and T3 and D50 recognized the trimer at greater levels than gp140(-). Yang et al. [2000] (vaccine antigen design)
- T4: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits - a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen - SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the

only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999] (vaccine antigen design)

- T4: This antibody, along with 7 others (M10, D41, D54, T6, T9, T10 and T35), can block the linear murine MAb D61, and the human MAb 246-D, which both bind to the immunodominant region near the two Cys in gp41 - most of these antibodies are oligomer dependent - all of the MAbs are reactive with ten different HIV-1 strains – members of this competition group are blocked by sera from HIV-1 + individuals. Earl et al. [1997] (antibody interactions)
- T4: MAbs T4 and T6 bind only to oligomer, and pulse chase experiments indicate that the epitope is very slow to form, requiring one to two hours. Otteken et al. [1996] (antibody binding site definition and exposure)
- T4: Does not bind to soluble monomeric gp41(21-166) that lacks the fusion peptide and membrane anchor, only to the oligomer gp140, as does T6. Weissenhorn et al. [1996] (antibody binding site definition and exposure)
- T4: one of five MAbs (T4, T6, T9, T10 and T35) in a competition group that bind to a conformation-dependent epitope in gp41 and is oligomer specific - neutralizes IIIB and SF2. Broder et al. [1994] (antibody binding site definition and exposure)
- T4: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994] (antibody generation)

No. 875

MAb ID m18 (M18, FAb M18)

**HXB2 Location** Env **Author Location** Env

**Epitope** 

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

Research Contact D. S. Dimitrov

References McCaffrey et al. 2004; Zhang et al. 2003 Keywords antibody binding site definition and expo-

sure, antibody generation, subtype comparisons, vaccine antigen design, variant crossrecognition or cross-neutralization

- m18: Called M18. Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The loss of the glycan within the V3 loop (GM299 V3) or adjacent to V3 in C2 (GM292 C2), left SF162 susceptible to neturalization by FAb M18, and the glycan mutants in C3 (GM329 C3), C4 (GM438 C4), or V5 (GM454 V5) became resistant to M18 neutralization. The M18 epitope is unknown. V3 glycans tended to sheild V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans sheilded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004] (antibody binding site definition and exposure, vaccine antigen design)
- m18: m18 was selected from a human Fab phage display library by a new method called sequential antigen panning (SAP), using a series of antigens to screen the library to pick

broadly cross-reactive isolates. The epitope of m18 is independent of CD4 binding. The phage display library was constructed using the combined bone marrow of three long term non-progressors with potent NAb activity in their sera. m18 bound to gp140s from primary isolates from clades A-F with nM affinities. The ability to block cell mediated fusion by m18 was compared to Fabs X5 and b12 for a clade A, CRF01 EA, G, and 6 clade B isolates, and the inhibitory activity of m18 was slightly lower but comparable to neutralizing Fabs b12 and X5. It also showed broad cross-neutralization; 11/15 pseudotyped Envs from primary isolates from clades A-F were inhibited in an IC50 assay at concentration less than or equal to 100 ug/ml; X5 was also tested and somewhat more potent, generally requiring lower concentrations and inhibiting 13/15 primary isolates. Zhang et al. [2003] (antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 876

MAb ID multiple Fabs

**HXB2 Location** Env **Author Location** gp120

> Epitope Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Burton et al. 1991

A panel of anti-gp120 Fabs was generated by antigen selection from a random combinatorial library prepared from bone marrow from an asymptomatic individual. Burton et al. [1991]

No. 877

MAb ID multiple MAbs

**HXB2 Location** Env **Author Location** gp120

Epitope

Neutralizing Immunogen vaccine

Vector/Type: protein HIV component:

gp120

Species (Isotype) mouse

References Denisova et al. 1996

• When gp120 was used as an immunogen, in contrast to gp120 bound to an anti-V3 MAb, few MAbs were generated and all bound better to the native than to the denatured protein – MAbs generated were: G1B12, G2F7, G9G8, G12F12, G1B8, G11F11, G9E8, G1B11, G1B6, G6F2, G2E7. Denisova *et al.* [1996]

**No.** 878

MAb ID multiple MAbs

**HXB2 Location** Env **Author Location** gp120

Epitope

Neutralizing

Immunogen vaccine

Vector/Type: gp120-CD4 complex HIV

component: gp120

Species (Isotype) mouse

References Denisova et al. 1996

When gp120-CD4 was used as an immunogen, in contrast to gp120 bound to an anti-V3 MAb, few MAbs were generated and all bound better to the native than to the denatured protein – MAbs generated were: CG43, CG41, CG49, CG53, CG42, CG4, CG46, CG40, CG52, CG51, CG48, CG50, CG125, CG124, CG121. Denisova et al. [1996]

No. 879

MAb ID multiple MAbs

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein-Ab complex HIV com-

ponent: gp120-Mab complex

Species (Isotype) mouse

References Denisova et al. 1996

• When anti-V3 MAb M77 was bound to gp120 and used as an immunogen, it stimulated many MAbs to linear epitopes, as well as an array of MAbs to discontinuous epitope – 10 of 36 MAbs were mapped to linear epitopes and are mentioned elsewhere in this database, the others are: GV5H1, GV4D5, GV4G10, GV1A8, GV10H5, GV8E11, GV2H4, GV6E6, GV1F7, GV1G9, GV4G5, GV6B12, GV1E8, GV2B7, GV1B11, GV6H5, GV6G2, GV6B5, GV1E10, GV5E3, GV5B9, GV5F4, GV6G4, GV1A12, GV5C11, GV6B6, GV3C10. Denisova *et al.* [1996]

No. 880

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Neutralizing LP

Immunogen HIV-1 infection

Species (Isotype) human (IgG3)

**References** Scharf *et al.* 2001

• IgG3: HIVIG was separated into immunoglobulin classes and IgG3 neutralization of HIV strains X4, R5 and X4R5 strains was superior to IgG1 and IgG2, and IgG3 was also a more potent inhibitor of viral fusion – the IgG3 advantage was lost when only Fabs were considered, indicating the IgG3 neutralization efficacy is enhanced due to a longer hinge region of the heavy chain in comparison to IgG1 and IgG2. Scharf *et al.* [2001]

No. 881

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp140 (IIIB)

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp120, gp140 Adjuvant:

MPL-SE adjuvant, QS21

Species (Isotype) rabbit (IgG)

References Earl et al. 2001

• Immunization of rabbits with oligomeric gp140 induced production of higher levels of cross-reactive neutralizing Abs than immunization with gp120 – immunization of Rhesus macaques with gp140 yielded strong NAb against IIIB, modest against other lab-adapted strains, and no NAb activity against primary isolates – most neutralizing activity could not be blocked by a V3 peptide – 3/4 vaccinated macaques showed no viral replication upon intravenous challenge with SHIV-HXB2. Earl *et al.* [2001]

**No.** 882

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp160 (IIIB)

**Epitope Neutralizing** 

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: gp160 Adjuvant: aluminum hydroxide

Species (Isotype) human

References Cox et al. 1999

• 60 asymptomatic HIV-1 infected patients were vaccinated with rec gp160 in alum, produced in a baculovirus expression vector in insect cells (VaxSyn), 64 received placebo, and all were followed in a 5 year longitudinal study – a mean of 78% of vaccinated and 82% of those receiving placebo had demonstrable ADCC at the different time intervals in the study, and the vaccine did not enhance ADCC production – patients with rapid and slow disease progression showed similar ADCC levels. Cox et al. [1999]

**No.** 883

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp160 (89.6)

Epitope Neutralizing yes Immunogen vaccine

Vector/Type: modified vaccinia Ankara (MVA) Strain: B clade 89.6 HIV component: Env, Gag-Pol Adjuvant: IL-2/Ig

Species (Isotype) macaque

References Barouch et al. 2001b

• Four rhesus macaques were vaccinated with a modified vaccinia Ankara (MVA) vaccine that elicited strong CTL responses as well as antibody responses. The animals were infected when challenged with pathogenic SHIV-89.6P, but had potent CTL responses, secondary NAb responses upon challenge, partial preservation of CD4+ T-cell counts, lower viral loads, and no evidence of disease or mortality by day 168 after challenge—monkeys that got a sham vaccine had high viral load, progressed to disease, and 2/4 were dead by day 168. Barouch et al. [2001b]

No. 884 MAb ID polyclonal HXB2 Location Env Author Location gp160 Epitope Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human

References Ahmad et al. 2001

 High CD4+ T-cell count and low viral load was correlated with high ADCC anti-HIV-1 Env Ab titers in a study of 46 HIV-1 infected individuals from all disease stages. Ahmad et al. [2001]

No. 885

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp160

**Epitope** 

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Beirnaert et al. 2001

• Neutralizing antibodies are thought to inhibit HIV entry by blocking either binding or fusion – six broadly cross-neutralizing sera that can neutralize group M and O viruses inhibit the binding to PBMCs – the nine primary isolates tested in this study represented very diverse subtypes and recombinant forms, and different co-receptor usage. Beirnaert *et al.* [2001]

**No.** 886

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp160

**Epitope** 

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Beirnaert et al. 2000

• Sera from 66 HIV individuals from diverse geographic locations could neutralize primary isolates to different extents: broad cross-neutralizing isolates could neutralize 14 primary isolates from HIV-1 group M clades A-H and three O isolates, limited cross-neutralizing sera neutralized some isolates, and non-neutralizing sera—6/7 broadly neutralizing sera were from African women, despite only 14/66 study subjects being women—ability to neutralize three key isolates, MN lab (envB/gagB, X4 coreceptor), VI525 (envG/gagH, envA/gagA, R5X4) and CA9 (Group O, R5) was predictive of being able to neutralize an additional set of 14 primary isolates. Beirnaert et al. [2000]

No. 887

MAb ID polyclonal

HXB2 Location Env

Author Location gp120 (SF2)

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade SF2 HIV component: gp120 Adjuvant: MF59,

PLG

Species (Isotype) mouse, baboon

References O'Hagan et al. 2000

Microparticles were used as an adjuvant for entrapped HIV-1 gp120 and induced strong serum IgG responses in mice – polylactide co-glycolide polymer (PLG) microparticles in combination with MF-59 had the highest response. O'Hagan *et al.* [2000]

No. 888

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (SF2, US4)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: DNA, protein Strain: B clade SF2, B clade US4 HIV component: gp120 Adjuvant: aluminum phosphate, MF59, PLG

Species (Isotype) macaque, guinea pig, mouse

References O'Hagan et al. 2001

• DNA vaccines of codon-optimized Env and Gag genes driven by CMV promotors and absorbed on to PLG microparticles were more effective than naked DNA at eliciting strong Ab responses (more rapid, higher titer, more stable), comparable to gp120 in MF-59. O'Hagan *et al.* [2001]

No. 889

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) chimpanzee (IgG)

References Moore & Burton 1999; Shibata et al. 1999

- polyclonal: Commentary discussing this finding noting the particularly high neutralization titer and limited breadth of the chimpanzee sera used in this study. Moore & Burton [1999]
- polyclonal: Purified IgG from chimpanzee sera infected with several HIV-1 strains was used for passive administration to macaques which were subsequently challenged with the virulent SHIV bearing the HIV-1 env DH12 *in vitro* neutralization correlated with protection *in vivo*. Shibata *et al.* [1999]

**No.** 890

MAb ID polyclonal

HXB2 Location Env

Author Location gp160 (MN)

**Epitope** 

**Neutralizing** LP

Immunogen HIV-1 infection

Species (Isotype) human (IgA)

References Moja et al. 2000

 15 samples isolated from parotid saliva were selected for study as they had anti-Env IgA – IgA neutralizing activity was detected that was not directed at either EDELKWA or the V3 loop. Moja et al. [2000]

No. 891

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

*Vector/Type:* protein *Strain:* B clade MN, B clade SF2 *HIV component:* gp120

Species (Isotype)

References McElrath et al. 2000

After 3 immunizations, 210/241 (87%) HIV-1 uninfected vaccinees in a phase II trial developed NAbs – of 140 patients receiving 4 vaccinations, 53% had persistent neutralizing antibodies to homologous virus, and 34% to heterologous virus, measured at day 728 after initial immunization – immunogens were well tolerated – but IVDUs had a decreased Ab response relative to lower risk groups. McElrath et al. [2000]

No. 892

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: gp120 Adjuvant: GM-CSF/gp120 chimera

Species (Isotype) mouse

References Rodríguez et al. 1999

The murine Ab response to a chimeric of granulocyte-macrophage colony stimulating factor GM-CSF/gp120 in vaccinia was not higher titer that the response to a gp120-vaccinia construct, but the breadth of the Ab response was greater – a cellular response of greater intensity was trigged to the GM-CSF/gp120 vaccinia construct, as measured by proliferation and Elispot. Rodríguez *et al.* [1999]

No. 893

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (YU2)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: stabilized Env trimer Strain: B clade HXBc2, B clade YU2 HIV component: Env

Species (Isotype) mouse (IgG)

Research Contact Joseph Sodroski, Harvard Medical School References Yang *et al.* 2001

• Soluble Env trimers were created that were designed to mimic functional Env oligomers – stabilized timers could induce neutralizing antibodies more effectively than gp120, and Abs to the YU2 trimer were cross-reactive within clade B and could neutralize several primary and TCLA reactive strains – the stabilized primers did not neutralize primary isolates outside the B clade, from clades C, D, and E – HXBc2 stabilized trimer antigen elicited strong neutralizing Abs against the homologous isolate HXBc2 TCLA strain, but not against primary isolates. Yang et al. [2001]

No. 894

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (MN)

**Epitope Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade MN HIV component: gp120 Adjuvant: aluminum hydroxide, QS21

Species (Isotype) human

References Evans et al. 2001

• Vaccination with QS21 adjuvant and rsgp120 elicited stronger and more sustained neutralizing antibody responses and lymphocyte proliferation with lower doses of rsgp120 than alum formulations, suggesting QS21 may be a means to reduce the does of soluble protein. Evans *et al.* [2001]

No. 895

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing yes

Immunogen HIV-1 infection

Species (Isotype) human

References Binley et al. 2000

HAART inhibited the development of anti-gp120 Ab when initiated during primary infection and sometimes in patients treated within 2 years of HIV-1 infection – HAART during primary infection usually did not inhibit the development of weak NAb responses against autologous virus – 3/4 patients intermittently adherent developed high titers of autologous NAbs, largely coincident with brief viremic periods. Binley et al. [2000]

No. 896

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (SIV)

**Epitope** 

Neutralizing yes

Immunogen HIV-1 infection

Species (Isotype) macaque

References Reitter et al. 1998

• This study concerned an SIV mutated strain that lacked 4th, 5th and 6th sites for N-linked glycosylation – monkeys infected with the mutant viruses had increased neutralizing activity in their sera relative to monkeys infected with the parental strain. Reitter *et al.* [1998]

**No.** 897

MAb ID polyclonal

**HXB2 Location** Env

Author Location Env

Epitope

**Neutralizing** yes

Immunogen HIV-1 infection

Species (Isotype) human

References Kim et al. 2001

After HAART reduction of viral load to <400 for three visits over a 12 month interval, 2/11 patients were found to have increased anti-Env Ab binding titers, and neutralizing Abs titers increased against primary isolates US1, and CM237 – no NAB titer increase was seen to more readily neutralized isolate BZ167 – this suggests that in certain individuals the control of HIV-1 by HAART may augment immune control of HIV. Kim et al. [2001]</li>

No. 898

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Neutralizing yes

Immunogen HIV-1 exposed seronegative

Species (Isotype) human (IgA)

**References** Kaul *et al.* 2001b

• Kaul *et al.* provide a concise summary of the findings concerning the presence of Mucosal IgA in highly exposed, uninfected subjects, arguing for a role in protection. Kaul *et al.* [2001b]

No. 899

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** yes

Immunogen vaccine

Vector/Type: protein Strain: B clade SF2 HIV component: gp120 Adjuvant: MF59

Species (Isotype) human

References Nitayaphan et al. 2000

A phase I/II trial was conducted in 52 seronegative Thais immunizing with rgp120 SF2 – the vaccine was safe and 39/40 developed NAb responses to the autologous SF2, while 22/40 were able to cross-neutralize the heterologous strain MN. Nitayaphan et al. [2000]

**No.** 900

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (SF2)

Epitope

**Neutralizing** yes

Immunogen vaccine

Vector/Type: protein Strain: B clade SF2 HIV component: gp120, p24 Gag Adjuvant: Immune stimulating complexes (ISCOM)

Species (Isotype) macaque

References Heeney et al. 1998a

The immune responses induced in Rhesus monkeys using two different immunization strategies was studied – one vaccine group was completely protected from challenge infection, the other vaccinees and controls became infected – protected animals had high titers of heterologous NAbs, and HIV-1-specific T helper responses – increases in RANTES, MIP 1 alpha and MIP 1 beta produced by circulating CD8+ T cells were also associated with protection. Heeney et al. [1998a]

No. 901

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

Epitope

Neutralizing

Immunogen vaccine

Vector/Type: peptide, protein Strain: B clade SF2, B clade SF33 HIV component: gp120 Adjuvant: Immune stimulating com-

plexes (ISCOM), MF59

Species (Isotype) macaque

References Verschoor et al. 1999

 Attempts were made to broaden immune responses induced in Rhesus monkeys by immunization of animals previously immunized that had resisted homologous challenge, with a second immunization with ISCOM-peptides or a boost with gp120 from SF33 – animals didn't survive a second challenge heterologous challenge virus SHIV(SF33) raising concerns about early antigenic sin. Verschoor et al. [1999]

No. 902

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

**Neutralizing** yes

Immunogen vaccine

Vector/Type: protein Strain: B clade SF2, CRF01 CM235 HIV component: gp120 Adjuvant: MF59

najavani.

Species (Isotype) baboon

References VanCott et al. 1999

• Immunization with rgp120 CM235 (CRF01) induced Abs capable of neutralizing TCLA subtype E (CRF01) and subtype B isolates, while rgp120SF2 induced Abs could only neutralize subtype B TCLA isolates – neither immunogen induced Abs capable of neutralizing primary HIV-1 isolates – both rgp120CM235 and rgp120SF2 induced Abs to regions within C1, V1/V2, V3, and C5, but unique responses were induced by rgp120CM235 to epitopes within C2, and by rgp120SF2 to multiple epitopes within C3, V4, and C4 – CM235 baboon sera bound 3- to 12-fold more strongly than the SF2 baboon sera to all subtype E gp120s while binding to subtype B gp120s (except SF2) were within two to threefold for the SF2 and CM235 baboon sera. VanCott et al. [1999]

**No.** 903

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp140 (SF162DeltaV2)

**Epitope Neutralizing** yes

Immunogen vaccine

Vector/Type: DNA with CMV promotor Strain: B clade SF162 HIV component:

gp140 Adjuvant: MF59

Species (Isotype) macaque, rabbit (IgG)

References Barnett et al. 2001

• SF162ΔV2 is a virus that has a 30 amino acids deletion in the V2 loop that does not abrogate its infectivity but renders it highly susceptible to neutralization—when incorporated into a codon-optimized DNA vaccine with a CMV promoter and delivered by gene gun, SF162ΔV2 gave higher neutralizing Ab titers against SF162 than did SF162 itself, and Abs that cross-neutralized non-homologous primary isolates were obtained only when SF162ΔV2, but not intact SF162, was used as the immunogen-Control MAbs 2F5 and 2G12 could neutralize all of the following primary isolates: 91US056(R5), 92US714(R5), 92US660(R5), 92HT593(R5X4), and BZ167(R5X4), while after the first protein boost, the sera from two SF162ΔV2 immunized macaques could neutralize 91US056(R5), 92US714(R5), 92US660(R5) and ADA(R5), but not 92HT593(R5X4) or 92US657(R5) the pattern of cross-recognition shifted after the second boost. Barnett et al. [2001]

**No.** 904

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Binley et al. 1997b

Retention of anti-Env antibodies and loss of anti-Gag antibodies during progression was studied, and suggested to be the result of the loss of T-cell help and the unique ability of Env to stimulate B cells even in a backdrop of declining CD4 cells, because of the ability of Env to bind to the CD4 molecule. Binley et al. [1997b]

No. 905

MAb ID polyclonal

HXB2 Location Env

Author Location gp120 (W61D)

Epitope

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade W61D

HIV component: gp120

Species (Isotype) human

References Beddows et al. 1999

rgp120 derived from a R5X4 subtype B virus, HIV-1 W61D, was used to vaccinate healthy volunteers and the resulting sera were compared with HIV-1 positive subjects – vaccinee sera had more potent responses to linear V1/V2 and V3 epitopes than did the sera from HIV-1 + individuals, but could only neutralize homologous or heterologous virus only after adaptation to T-cell lines – neutralization activity was lost after re-adaptation to growth in PBMCs – in contrast, sera from infected individuals could neutralize both PBMC and T-cell line adapted viruses. Beddows et al. [1999]

**No.** 906

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

*Vector/Type:* virus-like particle (VLP) *HIV component:* Gag, gp120, V3

Species (Isotype) macaque

References Wagner et al. 1998b

A VLP is a non-infectious virus-like particle self-assembled from HIV Pr55 gag – macaques were immunized with VLPs bound to either gp120 or V3+CD4 linear domains – Gag and Env specific CTL were stimulated in each case, and Ab response to gag and gp120 and was elicited, but the gp120 neutralizing response occurred only with whole gp120, not V3+CD4 – despite the CTL and Ab response, immunized macaques were infected by intervenous challenge with SHIV chimeric challenge stock. Wagner et al. [1998b]

No. 907

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope Neutralizing** 

Immunogen vaccine

Vector/Type: DNA HIV component: gp120,

gp160

Species (Isotype) mouse

References Shiver et al. 1997

 DNA vaccinations of BALBc mice with a gp120 or gp160 DNA vaccine elicited a strong T cell proliferative response with Th1-like secretion of gamma interferon and IL-2, with little or no IL-4, as well as antigen specific gp120 Abs. Shiver et al. [1997]

**No.** 908

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: DNA HIV component: Env, Gag, Pol, Vif Adjuvant: B7, IL-12

Species (Isotype) mouse

References Kim et al. 1997b

 A gag/pol, vif or CMN160 DNA vaccine, when delivered in conjunction with the plasmid encoding the co-stimulatory molecules B7 and IL-12, gave a dramatic increase in both the cytotoxic and proliferative responses in mice – the Ab response was detected by ELISA, but the CMN160 DNA vaccinated mice showed a neutralizing Ab response. Kim et al. [1997b]

**No.** 909

MAb ID polyclonal

HXB2 Location Env

Author Location gp120

Epitope

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Bradney et al. 1999

 Sera were taken from long term non-progressors and evidence for viral escape was noted – serum could neutralize earlier autologous isolates, but not contemporary isolates. Bradney et al. [1999]

No. 910

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing LP

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade SF2 HIV compo-

nent: Env, Gag

Species (Isotype) human

References Belshe et al. 1998

• NAbs were obtained by a HIV-1 gag/env in canary pox vaccination of eight volunteers after boosting with rgp120 against lab strains – 1/8 primary isolates was neutralized, BZ167. Belshe *et al.* [1998]

No. 911

MAb ID polyclonal

HXB2 Location Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Gag, gp120,

Protease Adjuvant: MF59

Species (Isotype) human

References Belshe et al. 2001; Belshe et al. 1998

A phase 2 trial was conducted in 435 volunteers with vCP201, a canary pox vector carrying gp120 (MN in vCP201, and SF2 in the boost), p55 (LAI) and protease (LAI), either alone or with a gp120 boost – NAbs against MN were obtained in 56% of those who received vCP201 alone, and in 94% of those who got the prime with the gp120 boost. Belshe *et al.* [1998]

**No.** 912

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp120

Epitope

Neutralizing

Immunogen

Species (Isotype) human

**References** Neshat *et al.* 2000

• HIV-1 gp120 appears to be a B cell superantigen that binds to members of the  $V_{\rm H3}$  Ig gene family—the gp120 binding site was localized to the Fab portion of the Ab, and discontinuous residues in the  $V_{\rm H}$  region were critical. Neshat *et al.* [2000]

**No.** 913

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp41 (539-684 BH10)

Epitope Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: gp41

**Species (Isotype)** mouse (IgG) **References** Bai *et al.* 2000

• Murine rsgp41 antisera recognized a common epitope on human IFN $\alpha$  (aa 29-35 and aa 123-140) and on human IFN $\beta$  (aa 31-37 and aa 125-142), suggesting that elevated levels of Ab to IFNs found in HIV+ individuals may be due to a cross-reactive gp41 response. Bai *et al.* [2000]

No. 914

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (BH10)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: DNA Strain: B clade 89.6, B clade ADA, B clade IIIB HIV component:

gp120 Adjuvant: C3d fusion

**Species (Isotype)** mouse (IgG) **References** Ross *et al.* 2001

gp120 was fused with murine complement protein C3d in a DNA vaccine to enhance the titers of Ab to Env – fusion to C3d resulted in a more rapid onset of Ab response and avidity maturation, after three immunizations in BALB/c mice with DNA on a gold bead delivered with a gene gun, but not in strong neutralizing Ab response. Ross et al. [2001]

No. 915

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (SF162DeltaV2)

Epitope Neutralizing

Immunogen vaccine

Vector/Type: DNA prime with protein boost Strain: B clade SF162 HIV component:

gp140 Adjuvant: MF59

Species (Isotype) macaque

**References** Cherpelis *et al.* 2001a; Cherpelis *et al.* 2001b

- Two animals were immunized both intradermally and intramuscularly at weeks 0, 4, and 8 with a codon optimized DNA vector expressing the SF162V2 gp140 envelope with an intact gp120-gp41 cleavage site, and both developed lymphoproliferative responses and potent neutralizing Abs CD8+ T lymphocytes were depleted in the animals and they were challenged with SHIV162P4 at peak viremia, plasma viral levels in the vaccinated animals were 1 to 4 logs lower than those in the unvaccinated animals. Cherpelis *et al.* [2001b]
- HIV-1 SF162ΔV2 gp140 envelope was used in a DNA-prime plus protein-boost vaccination methodology in Rhesus macaques, the animals were depleted of their CD8+ T lymphocytes, and challenged with pathogenic SHIV(SF162P4)—the vaccinated macaques had lower peak viremia, rapidly cleared

virus from the periphery, and developed delayed seroconversion to SIV core antigens relative to non-vaccinated controls. Cherpelis *et al.* [2001a]

No. 916

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Sarmati et al. 2001

 Some HIV-1 infected patients have increasing CD4 counts despite failing ARV, and CD4 levels are correlated with HIV-1 specific NAbs – no correlation was found between NAbs and viral load in this patients. Sarmati et al. [2001]

No. 917

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp41 (539–684 BH10)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: gp41

Species (Isotype) mouse (IgG)

References Bai et al. 2000

 There is a common epitope in HIV-1 gp41, and IFNalpha and IFNbeta. Bai et al. [2000]

**No.** 918

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype) human (IgM)

References Llorente et al. 1999

 Combinatorial antibody analysis by phage display and flow cytometry demonstrated that gp120 in HIV-1 negative people is recognized by IgM, but not IgG Abs – IgM Fab reactivity is observed throughout the entire sequence of HIV-1 IIIB gp120 and is characterized by low affinity binding and near germline configuration reflecting a lack of maturation of the IgM repertoire – no neutralizing activity was observed in a non-infected individual before isotope switching. Llorente *et al.* [1999]

No. 919

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp120 (SF2)

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade SF2

HIV component: gp120

Species (Isotype) human (IgM)

References Locher et al. 1999

High risk volunteers were vaccinated with SF2 gp120 – 3 breakthrough cases were studied – SF2 neutralizing Abs were observed, but Ab titers to autologous virus were never high and took 6 months after HIV-1 infection to develop – viral loads were similar to HIV-1 infected individuals who had not been vaccinated. Locher et al. [1999]

**No.** 920

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (subtype A, B, C, D, CRF01)

**Epitope** 

Subtype A, B, C

Neutralizing yes

Immunogen vaccine

*Vector/Type:* formaldehyde-fixed whole-cell

HIV component: gp120

Species (Isotype) mouse (IgG)

References Nunberg 2002; LaCasse et al. 1999

- A retraction was printed (Science 296:1025, 2002) noting that an unknown cytotoxic effect of these complex sera accounted for a major fraction of the neutralization reported in LaCasse *et al.* [1999] Nunberg [2002]. LaCasse *et al.* [1999]; Nunberg [2002]
- In this study, immunogens were generated that were thought to capture transient envelope-CD4-coreceptor structures that arise during HIV binding and fusion by formaldehyde-fixation of co-cultures of cells expressing HIV-1 Env and those expressing CD4 and CCR5 receptors – these cells elicited NAbs in CD4and CCR5-transgenic mice that neutralized 23/24 primary isolates from clades A-E. LaCasse et al. [1999]

**No.** 921

MAb ID polyclonal

HXB2 Location Env

Author Location (B consensus)

**Epitope** 

Subtype B

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Morris et al. 2001

Ab responses before HAART therapy and after one year of therapy were measured in 8 individuals that were classified HAART successes, and 10 patients who were classified as HAART failures – V3 peptide antibody binding titers to the B-consensus and MN and SF2 variants, and neutralization of HIV-1 MN and four subtype B clinical isolates were tested – subjects with strong anti-V3 and NAb humoral immune responses before starting HAART were more likely to achieve sustained viral suppression to <500 copies RNA/ml on HAART – HIV-specific Ab responses declined after 1 year of successful viral suppression on HAART. Morris et al. [2001]</li>

No. 922

MAb ID polyclonal

HXB2 Location Env

**Author Location** 

Epitope

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Pilgrim et al. 1997

• Sera from long-term nonprogressors(LTNP) had broader NAbs against heterologous primary isolates and were more likely to neutralize the contemporaneous autologous isolate than were sera from short-term nonprogressors and normal progressors — in 4 individuals followed from acute infection, NAbs were detected against the early autologous isolate by 5-40 weeks, and not detected in an additional 2 cases after 27-45 weeks. Pilgrim *et al.* [1997]

No. 923

MAb ID polyclonal

HXB2 Location Env

**Author Location** 

**Epitope** 

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Moog et al. 1997

 Autologous and heterologous NAbs were studied in 18 individuals who were sampled early after sero-conversion and followed longitudinally – autologous NAbs were not detected in sera collected at the same time as the viruses were isolated – NAbs detected against the seroconversion autologous strains were not detected one year after seroconversion, and were highly specific to the virus present at the early phase of HIV infection – heterologous neutralization of primary isolates were not detected until after 2 years. Moog et al. [1997]

No. 924

MAb ID polyclonal

HXB2 Location Env

**Author Location** 

Epitope

**Neutralizing** yes

Immunogen HIV-1 infection

Species (Isotype) human

References Montefiori et al. 2001

• In 7/9 patients in whom HAART was initiated during early seroconversion, NAbs to autologous strains were not found immediately following treatment interuption after 1-3 years, and Env and Gag Abs were low or undetected by ELISA indicating, that early HAART suppresses the normal antibody response to HIV-1, presumably by limiting the concentration of viral antigens needed to drive B-cell maturation – in 3 patients with a viral rebound autologous NAbs rapidly appeared and correlated with spontaneous down-regulation of viremia - prolonged control of viremia after stopping treatment persisted in the absence of detectable NAbs, suggesting that cellular immune responses alone can control viremia under certain circumstances - these results support the notion that virus-specific B-cell priming, combined with CD8+ CTL induction, may be beneficial for HIV-1 vaccines that aim to suppress viremia in the absence of complete protection to prevent disease and reduce the rate of virus transmission. Montefiori et al. [2001]

No. 925

MAb ID polyclonal

**HXB2 Location** Env **Author Location Epitope** Subtype B **Neutralizing** 

**Immunogen** HIV-1 infection Species (Isotype) human (IgG)

References Scala et al. 1999

• Random peptide libraries were screened using sera from HIVinfected subjects to identify mimotopes, peptides that mimic conformational or linear epitopes specifically recognized by Abs from HIV-1 infected individuals - the sera of simian SHIV-infected monkeys also recognized the specific peptides, and mice immunized with the selected peptides elicited HIVspecific Abs that neutralized HIV-1 isolates IIIB and NL4-3. Scala et al. [1999]

No. 926

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide HIV component: mimotopes

Species (Isotype) mouse (IgG)

References Scala et al. 1999

· Random peptide libraries were screened using sera from HIVinfected subjects to identify mimotopes, peptides that mimic conformational or linear epitopes specifically recognized by Abs from HIV-1 infected individuals - the sera of simian SHIV-infected monkeys also recognized the specific peptides, and mice immunized with the selected peptides elicited HIVspecific Abs that neutralized HIV-1 isolates IIIB and NL4-3. Scala et al. [1999]

No. 927

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: virus-like particle (VLP) HIV component: Env, Gag Adjuvant: Complete

Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgG)

References Lebedev et al. 2000

• Virus-like particles (VLPs) in the form of spherical particles with yeast dsRNA enveloped in a polysaccharide matrix carrying the protein TBI, that contains fragments of HIV Env and Gag, were used to immunize BALB/c mice and induced specific Abs against HIV-1 as measured by ELISA with TBI. Lebedev et al. [2000]

No. 928

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Donners et al. 2002

• A difference in neutralization patterns between African and European plasma is observed, especially in African women, who tended to have cross-neutralizing Abs against primary isolates. Donners et al. [2002]

No. 929

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Dianzani et al. 2002

• Immune complexes(ICs) in the plasma contained HIV RNA (80%-100%) in association with HIV-specific IgG NAbs indicating that the HIV in the plasma of carriers is frequently composed of antibody-neutralized HIV as ICs. Dianzani et al. [2002]

No. 930

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Neutralizing P

**Immunogen** HIV-1 infection

Species (Isotype) human (IgG)

References Kimura et al. 2002

Significant neutralization activity against autologous isolates was observed in 13/19 HIV+ patients at initiation of HAART therapy which persisted during therapy, increasing in one patient, and declining in one patient – 3/6 patients with no detectable NAb at the start of therapy developed NAb responses – of the four patients with increased NAb responses, three had low level viral rebounds (blips). Kimura et al. [2002]

No. 931

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing P

Immunogen HIV-1 exposed seronegative

Species (Isotype) human (IgA)

References Devito et al. 2000b

· Mucosal and plasma HIV-specific IgA that can neutralize primary isolates is present saliva (11/15 tested) and plasma (11/15) and cervicovaginal fluid (11/14) from highly exposed persistently seronegative (HEPS) individuals. Devito et al. [2000b]

No. 932

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing P

Immunogen HIV-1 exposed seronegative

Species (Isotype) human (IgA)

References Devito et al. 2000a

IgA from the genital tract, saliva and plasma from highly exposed persistently seronegative (HEPS) individuals can inhibit transcytosis of HIV-1 across a transwall system that provides a tight epithelial cell layer—50% of the IgA samples studied were able to inhibit transcytosis of at least one of two primary isolates tested, indicating this may be an important mechanism against sexual acquisition of HIV-1. Devito et al. [2000a]

No. 933

MAb ID polyclonal

HXB2 Location Env

**Author Location** 

**Epitope** 

Subtype A, B, D

Neutralizing P

Immunogen HIV-1 exposed seronegative

Species (Isotype) human (IgA)

References Broliden et al. 2001

 IgA isolated from the saliva, genital tract, and plasma of most highly exposed persistently seronegative (HEPS) sex workers in a Kenyan cohort could neutralize a B, A and D clade primary isolates and could inhibit transcytosis of HIV across a transwall model of the human mucosal epithelium. Broliden et al. [2001]

**No.** 934

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype A, B, D

**Neutralizing** P

Immunogen HIV-1 exposed seronegative

Species (Isotype) human (IgA)

References Devito et al. 2002

• IgA isolated from the saliva, genital tract, and plasma of most highly exposed persistently seronegative (HEPS) Kenyan sex workers mediated broad cross-clade neutralization of primary isolates (A, B, C, D, and CRF01) – 6/10 HEPS individuals that were persistently exposed to a stable HIV+ B clade infected partner showed less breadth of neutralization, and were able to neutralize clade A and B primary isolates, but not clades C, D, or CRF01. Devito *et al.* [2002]

**No.** 935

 $\boldsymbol{MAb\;ID}\;\; polyclonal$ 

HXB2 Location Env

**Author Location** 

**Epitope** 

Neutralizing P

Immunogen HIV-1 exposed seronegative

Species (Isotype) human (IgA)

References Mazzoli et al. 1999

Serum HIV-specific IgA is present in highly exposed persistently seronegative individuals (HEPS) in the absence of serum IgG – serum IgA can be found in productively infected individuals and exposed seronegatives at similar titers – 5/15 sera from HEPS had neutralizing activity, 2 of these in purified IgA – HIV-1 specific serum IgA concentrations declined after one year of interruption of at-risk sex. Mazzoli *et al.* [1999]

No. 936

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing P

Immunogen HIV-1 exposed seronegative

Species (Isotype) human (IgA)

References Beyrer et al. 1999

 HIV-specific anti-gp160 IgA is present in cervical lavage from 6/13 HIV-exposed seronegative Thai female sex workers. Beyrer et al. [1999]

No. 937

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: DNA Strain: B clade

HXB2/Bal

Species (Isotype) mouse

References Chakrabarti et al. 2002

• A modified gp140 (gp140ΔCFI), with C-term mutations intended to mimic a fusion intermediate and stabilize trimer formation, retained antigenic conformational determinants as defined by binding to CD4 and to MAbs 2F5, 2G12, F105, and b12, and enhanced humoral immunity without diminishing the CTL response in mice injected with a DNA vaccine. Chakrabarti *et al.* [2002]

**No.** 938

MAb ID polyclonal

HXB2 Location Env

**Author Location** 

**Epitope** 

Subtype C

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6, B clade IIIB HIV component: Env Adjuvant: alpha2-macroglobin, Complete Freund's Adjuvant (CFA), GM-CSF, monophosphoryl

lipid A

Species (Isotype) mouse

References Liao et al. 2002

 HIV-envelope peptides coupled to α2-macroglobin were much more immunogenic when formulated in monophosphoryl lipid A with GM-CSF than in complete or incomplete Freund's adjuvant or in monophosphoryl lipid A with GM-CSF alone. Liao *et al.* [2002]

No. 939

MAb ID polyclonal

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing P

Immunogen vaccine

Vector/Type: gp120-CD4 complex, gp140-CD4 complex Strain: B clade IIIB HIV component: gp120, gp140 Adjuvant: QS21

Species (Isotype) macaque

References Fouts et al. 2002

• gp120-CD4 and gp140-CD4 complexes were used for i.m. vaccination of rhesus macaques and neutralizing Ig was recovered using affinity chromatography using a chimeric HIV-BAL gp120 with a mimetic peptide that induces a CD4-triggered mimetic structure – the sera and affinity purified Ab were broadly neutralizing against primary X4, R5, and R5X4 isolates from multiple subtypes but did not react as well against lab-adapted isolates. Fouts *et al.* [2002]

**No.** 940

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Pastori et al. 2002

• HAART initiated during primary infection was studied in seven patients and had different effects on NAb production—in some cases, α-Env Abs were inhibited during primary infection, and in some cases strong NAbs against autologous virus were induced. Pastori *et al.* [2002]

**No.** 941

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen HIV-1 infection

Species (Isotype) chimpanzee (IgG)

References Moore & Burton 1999; Igarashi et al. 1999

• The rate of virus clearance in the circulation in rhesus macaques receiving a continuous infusion of cell-free viral dual-tropic virus isolate HIV-1DH12 particles in the presence and absence of virus-specific antibodies was measured – the clearance of physical and infectious viral particles is very rapid in naive animals, with half-lives ranging from 13 to 26 minutes, but clearance cold be acheived with a half life of 3.9-7.2 minutes when chimpanzee neutralizing Abs were present to help to remove virions from the blood. Igarashi *et al.* [1999]

 polyclonal: Commentary discussing this finding noting the particularly high neutralization titer and limited breadth of the chimpanzee sera used in this study. Moore & Burton [1999]

No. 942

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: gp120, gp41

Adjuvant: MF59

Species (Isotype) human

References Gupta et al. 2002

- Different HIV strains were used for different regions: gp120 MN and gp41 LAI, rgp120 SF2.
- Vaccine trial protocol 022A in 150 HIV-1 uninfected adults (130 completed the study) showed high titer ALVAC vaccine in combination with gp120 was safe and immunogenic in HIV-1 negative volunteers – NAb responses were detected in 95% of vaccinees, with higher titers in recipients of sequential versus simultaneous dosing of the two vaccines and in vaccinia naive volunteers. Gupta et al. [2002]

No. 943

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Neutralizing yes

Immunogen vaccine

Vector/Type: protein Strain: B clade 89.6 HIV component: gp120, gp140 Adjuvant:

Cholera toxin (CT), IL-12

Species (Isotype) mouse (IgA, IgG, IgG1, IgG2a)

References Albu et al. 2003

Keywords genital and mucosal immunity, mucosal im-

munity, Th1, Th2

 Mice were intranasally immunized with gp120 or gp140 with IL-12 and Cholera toxin as adjuvants. Adjuvants enhanced NAb stimulation in mucosa and genital tissues and in serum. Albu et al. [2003] (genital and mucosal immunity, mucosal immunity, Th1, Th2)

No. 944

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Subtype A

Neutralizing yes

Immunogen vaccine

Vector/Type: virus-like particle (VLP) Strain: A clade UG5.94UG018 HIV com-

ponent: Gag, gp120

Species (Isotype) mouse

References Buonaguro et al. 2002

Country Uganda

**Keywords** subtype comparisons

BALB/c mice were immunized with VLPs carrying a subtype A gp120. Humoral immune responses directed against B-clade derived Gag (p24) peptides or gp120-Env V3 loop peptide were readily induced following a multi-dose immunization with VLP particles presenting a gp120 molecule from a HIV-1 isolate of clade A. VLP-immunized mice showed autologous and heterologous (against B-clade HIV-1 IIIB strain) neutralization activity. Proliferative responses and CTL were also observed. Buonaguro et al. [2002] (subtype comparisons)

No. 945

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Subtype A, B, D

Neutralizing

Immunogen vaccine

Vector/Type: canarypox, protein Strain: B clade LAI, B clade MN HIV component: Env, Gag, Protease

Species (Isotype) human

References Cao et al. 2003

Country Uganda

**Keywords** subtype comparisons

• 20 Ugandan seronegative individuals were intramuscularly immunized in this study with an ALVAC HIV GagPol and Env vaccine carrying B clade agtigens.3/20 of subjects produced neutralizing antibodies against the autologous HIV-1 clade B strain MN that was T-cell line adapted; 2 also had NAb reactivity against a primary B clade cell line. No NAb cross-reaction was observed with primary viral isolates UG92029 (subtype A) or UG92046 (subtype D). 4/20 had detectable CTL activity against B clade antigen, and one of these cross-reacted with A clade antigen, one with D clade. Cao et al. [2003] (subtype comparisons)

**No.** 946

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp160

**Epitope** 

**Neutralizing** 

Immunogen SHIV infection

Species (Isotype) macaque

References Crawford et al. 1999

Keywords variant cross-recognition or crossneutralization

Neutralization assays with rsCD4, MAbs, and serum samples from SHIV-infected macaques and HIV-1 infected individuals were used to characterize the antigenic properties of the env glycoprotein of six primary isolate or TCLA SHIV variants. SHIV infected macaques could neutralize autologous virus very effectively, but serum from HXB2c or 89.6 infected animals could not neutralize heterologous SHIVs. Serum from KU infected animals could neutralize only HXB2c, and serum from 89.6PD infected animals could neutralize 89.6, 89.6P, 89.6PD and KB9 (all derived from 89.6) well. Many sera

from the SHIV infected macaques could also neutralize HIV-1 strains MN and SF2. Crawford *et al.* [1999] (variant cross-recognition or cross-neutralization)

No. 947

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp160

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Crawford et al. 1999

Keywords variant cross-recognition or crossneutralization

• Neutralization assays with rsCD4, MAbs, and serum samples from SHIV-infected macaques and HIV-1 infected individuals were used to characterize the antigenic properties of the env glycoprotein of six primary isolate or TCLA SHIV variants. Serum from 9 HIV-1 infected people were tested for their ability to neutralize SHIVs. KU2 was least sensitive, 89.6, 89.6P, 89.6PD and KB9 (all derived from 89.6) were moderately susceptible, and SHIV HXB2c was less sensitive than IIIB, the strain from which it was derived. Crawford et al. [1999] (variant cross-recognition or cross-neutralization)

No. 948

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Subtype B, C, CRF01\_AE

Neutralizing yes

Immunogen vaccine

Vector/Type: Venezuelan equine encephalitis virus (VEE) Strain: B clade R2 HIV com-

ponent: gp160∆CT

Species (Isotype) rabbit, mouse (IgG)

References Dong et al. 2003

**Keywords** subtype comparisons, variant cross-recognition or cross-neutralization

- Subcutaneous or intradermal immunization with VEE replicons expressing HIV-1 R2 gp140 and with HIV-1 R2 gp160 lacking the cytoplasmic tail. Sera from 3/3 rabbits inhibited SF162 infectivity and 2/3 rabbits were able to neutralize the R2 strain.
- C3H/He mice immunized with replicons expressing RT env protein or the VEE env vector pGP expressing either gp140 or gp160 showed cross-reactive neutralizing Ab responses to five clade B env proteins, a chinese clade C strain and weakly against a chinese clade E (CRF-1) strain.
- Mice and rabbits were immunized with Venezuelan equine encephalitis virus (VEE) replicon system particles expressing HIV-1 Env from the clone R2 that was derived from a virus that was neutralization sensitive and isolated from an individual that made strong NAb responses. Stronger and faster NAb responses were induced with replicons expressing gp160 with the cytoplasmic tail deleted than with gp160 or gp140. NAb responses against heterologous strain SF162 were similar in BALB/c and C3H/He mice and enhanced compared to

responses elicited in C57BL/6 mice. Serum from mice neutralized 5 primary clade B env proteins, a chinese clade C strain. but not a chinese clade E (CRF-1) strain. Sera from 3/3 immunized rabbits could neutralize SF162, and from 2/3 neutralized the autologous R2 strain. Dong et al. [2003] (variant crossrecognition or cross-neutralization, subtype comparisons)

No. 949

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype multiple, M, O

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Donners et al. 2003

Country Belgium

Keywords assay development, assay standardization/improvement, co-receptor, subtype comparisons

• Plasma samples from six HIV-1 + Belgians showed broad crossneutralization ability against primary isolates from group M (subtypes A-H) and Group O. Viruses with R5, X4, and R5X4 co-receptor usage were all represented in the test panel. Kinetics of neutralization showed that NAb responses detected using a PBMC assay with a short incubation period could be lost upon extended culture. No preincubation with Ab was needed to see some inhibition of virus replication, indicating that at least partial neutralization occurs post-virus binding to target cells. Donners et al. [2003] (assay development, coreceptor, kinetics, subtype comparisons, assay standardization/improvement)

No. 950

MAb ID polyclonal

**HXB2 Location** Env

Author Location Env

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Research Contact Rebeca Geffin, Miami School of Medicine

References Geffin et al. 2003

Keywords autologous responses, escape, rate of progression, responses in children

• A longitudinal study of NAb responses in perinatally HIV-1 infected infants and children was undertaken, including 7 with rapid progression (RP) and 9 who did not progress rapidly (NRP). A subset of both RPs and NRPs had some plasma samples that could neutralize contemporaneous autologous viral isolates after 6 months of age, but most isolates could not be neutralized by contemporaneous plasma, only by later samples. The non-contemporaneous NAbs would persist for years, had highest titers against earlier isolates, and tended to be more potent in NRP children. This study indicates that there is ongoing NAb escape in HIV-1 + children. No correlation between HIV RNA levels and Ab production was established, although this might have been complicated by treatment. Geffin et al. [2003]

(autologous responses, escape, responses in children, rate of progression)

No. 951

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype)

Research Contact Mascola2003b

References Mascola & Montefiori 2003

Keywords escape, review

• This paper reviews the paper by Wei et al. (Nature 2003) that substantiates the notion that HIV evolves to change the number and position of glycosylation sites in Envelope and this facilitates neutralization escape in vivo. This NAb escape mechanism is called a glycan shield. Mascola & Montefiori [2003] (escape, review)

No. 952

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

**Neutralizing** 

**Immunogen** 

Species (Isotype) macaque

References Mascola 2003

**Keywords** immunoprophylaxis, review

• This review dicusses the importance and function of protective antibody responses in animal model studies in the context of effective vaccine development. SHIV models have shown protection using high levels of MAbs can prevent infection, and partial protection that can influence disease course can be obtained from modest levels of NAbs. The binding properties and SHIV challenges studies conducted with infusions of combinations of MAbs b12, 2G12, and 2F5 are reviewed. Mascola [2003] (immunoprophylaxis, review)

No. 953

MAb ID polyclonal

HXB2 Location Env

Author Location gp120

**Epitope** 

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: DNA prime with virus-like particle (VLP) boost, fowlpoxvirus prime with virus-like particle (VLP) boost Strain: B

clade 89.6P HIV component: Env

Species (Isotype) rabbit

References Radaelli et al. 2003

Keywords Th1, Th2

• Three different immunization protocols using two recombinant fowlpox (FP) constructs and two expression plasmids (SIV mac239 gg/pol or HIV-1 env 89.6P) for priming and VLP particles for boosting were tested for their ability to elicit neutralizing Ab and cell-mediated immune responses. NAb responses against SHIV 89.6P were elicited in all protocols tested. Plasmid DNA (pcDNA3gag/pl SIV) was more efficient than the FP vector (FPgag/polSIV) in inducing Ab responses to the gag core protein (p27). DNA plasmid followed by a VLP boost elicited a Th0 profile. Radaelli *et al.* [2003] (Th1, Th2)

No. 954

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype B, CRF01\_AE

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Polonis et al. 2003

Country Thailand

Keywords co-receptor, escape, subtype comparisons

• Neutralization of 49 subtype E HIV-1 isolates from various stages of disease and 21 subtype B viruses was compared using polyclonal Ab pools and single subtype E plasmas. Non-syncytium-inducing (NSI) CRF01 (subtype E) HIV-1 isolates showed increased sensitivity to neutralization (42%) than syncytium-inducing (SI) subtype E isolates (9%). In contrast, the viral phenotype of subtype B isolates did not correlate with neutralizaiton sensitivity. SI viruses were primarily X4 (one X4R5 was identified), NSI were R5. Low CD4+ T cell numbers in subtype E infected patients correlated with concurrent isolate resistance to neutralizing Ab responses. Polonis *et al.* [2003] (co-receptor, escape, subtype comparisons)

No. 955

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade W61D HIV component: gp120, Nef, Tat Adjuvant: AS02A (oil-in-water emulsion, 3D-monophosphoryl lipid A, QS21), AS06 (CpG, aluminum hydroxide)

Species (Isotype) macaque (IgG)

References Voss et al. 2003

**Keywords** adjuvant comparison, variant crossrecognition or cross-neutralization

 Monkeys were immunized with different combinations of gp120/NefTat/SIV-Tat three times over a three-month time period and intravenously challenged with SHIV 89.6P four weeks after the last immunzation. Vaccination induced anti-Tat, -Nef and -gp120 Ab responses that decreased until challenge. Neutralizing Ab responses against HIV-1 MN and HIV-1 W61D were observed. Vaccination with the combination of all three proteins reduced viral load and kept animals from a CD4+ T cell decline, preventing AIDS for more than 2.5 years. The adjuvant AS02A elicited NAbs and protected from disease, while AS06 did not. gp120 alone or TatNef/SIV-Tat without gp120 could not protect from disease. Voss *et al.* [2003] (adjuvant comparison, variant cross-recognition or cross-neutralization)

No. 956

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: peptide Adjuvant: QS21

Species (Isotype) mouse

References Cunto-Amesty et al. 2001

Keywords mimotopes, vaccine antigen design

• Concanavalin A binds to mannose/glucose, and binds to HIV-1. Con A was used to select peptide mimics of carbohydrates that bound to Con A, and the mimetic peptides were then used for BALB/c mouse immunization. Abs raised against the mimetic peptides binds to HIV+ cells, and could weakly neutralize T cell lab adapted strains. Cunto-Amesty *et al.* [2001] (mimotopes, vaccine antigen design)

No. 957

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

Epitope

**Neutralizing** 

Immunogen vaccine

Vector/Type: E. Coli recombinant protein HIV component: gp120, gp41 Adjuvant: Complete Freund's Adjuvant (CFA), Incomplete Freund's Adjuvant (IFA)

Species (Isotype) mouse

References Li et al. 2002

**Keywords** vaccine antigen design

 A polyepitope vaccine was designed based on a recombinant GST fusion protein containing three repeats of the 2F5 core epitope ELDKWA combined with the V3 region peptide GP-GRAFY. Abs raised in mice could recognize the peptides, sgp41, and CHO-WT cells that expressed HIV-1 Env on their surface. Li et al. [2002] (vaccine antigen design)

No. 958

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Montefiori et al. 2003

**Keywords** acute/early infection, autologous responses, escape

• AC10 is a subject who was given treatment early after infection, and had a viral rebound after cessation of therapy, which then declined to a low level. The polyclonal sera from AC10 could potently neutralize the rebound virus, and NAb escape followed with a neutralizing response against the escape variant and subsequent escape from that response. Viral loads remained low in this subject despite escape. The rebound isolate that was potently neutralized by autologous sera was not particularly neutralization sensitive, as it resisted neutralization by sCD4 and MAbs IgG1b12, 2G12 and 2F5, and was only moderately sensitive to sera from other HIV+ individuals that had high titers of NAbs to TCLA strains. Montefiori et al. [2003] (autologous responses, acute/early infection, escape)

No. 959

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp120 (DH012)

Epitope Neutralizing

Immunogen HIV-1 infection, vaccine

Vector/Type: protein

**Species (Isotype)** chimpanzee **References** Zhu *et al.* 2003

Keywords vaccine-specific epitope characteristics

• This study compares the immunogenicity of the HIV DH012 strain in chimpanzees during a natural infection with DH012 vaccinations. Naturally infected chimpanzees have sera containing potent anti-DH012 neutralization Abs, but the primary epitope is a discontinuous conformational epitope called CEV that involves the V1/V2 region, the bridging sheet, and the V3 loop. Abs that are raised upon gp120 vaccination, in contrast, are primarily against V3. DH012 has preserved the epitopes for the MAbs IgG1b12, 2G12, 17b, however natural DH012 infection in chimpanzees and DH012 gp120 vaccination in guinea pigs does not give rise to Abs against these epitopes. Zhu et al. [2003] (vaccine-specific epitope characteristics)

**No.** 960

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp120 (IIIB)

**Epitope** 

Neutralizing yes

Immunogen HIV-1 infection

Species (Isotype) human

References Aasa-Chapman et al. 2004

**Keywords** acute/early infection, autologous responses

• Neutralizing Ab responses to autologous virus envelopes were studied in four acutely HIV-1 infected, treatment-naive, homosexual men (MM1,MM2, MM4 and MM8). Detection of gp120 antibodies was rapid using ELISPOT, within a few weeks, but detection of neutralizing antibodies took between 3 and 16 months, precluding involvement of detectable NAbs with resolution of viremia. Heterologous NAb responses arose even later, by 3 months or more, suggesting gradual broadening of the immune response. Aasa-Chapman et al. [2004] (autologous responses, acute/early infection)

**No.** 961

MAb ID polyclonal

**HXB2** Location Env

Author Location gp120 (V3) (IIIB)

Epitope Subtype B Neutralizing yes Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB, B clade MN HIV component: gp120 Adjuvant: Complete Freund's Adjuvant (CFA),

Incomplete Freund's Adjuvant (IFA)

Species (Isotype) rabbit, guinea pig

References Berman et al. 1992

Keywords vaccine-specific epitope characteristics

Abs derived from immunizations of rabbits and guinea pigs with either IIIB- or MN-gp120 were compared. Both could block gp120 binding to CD4, and this activity was strain-specific. Antisera from IIIB-rgp120 immunizations could only neutralize displayed homologous virus, while sera from MN-rgp120 rabbit vaccinations could neutralize MN 3/8 additional tested viruses. Berman *et al.* [1992] (vaccine-specific epitope characteristics)

No. 962

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env (YU-2)

Epitope
Subtype B
Neutralizing yes
Immunogen vaccine

Vector/Type: DNA with CMV promotor, DNA prime with protein boost Strain: B clade YU2 HIV component: gp140 Adjuvant: monophosphoryl lipid A, trehalose dicoryno-

mycolate

Species (Isotype) mouse (IgG)

References Bower et al. 2004

**Keywords** adjuvant comparison, vaccine antigen design A vaccines encoding an uncleaved form of YU-2 an140

• DNA vaccines encoding an uncleaved form of YU-2 gp140 stabilized with a synthetic trimerization domain isolated from the fibritin (FT) protein of the T4 bacteriophage and fused to murine C3d as a molecular adjuvant, could induce low titers of neutralizing antibodies against primary isolates HIV-1 YU-2 and HIV-1 ADA. DNA was administered by gene gun immunization to BALB/c mice, protein boost was performed by intraperitoneal injection. C3d is a component of the innate immune system that can serve as a moelcular adjuvant and had been previously shown to enhance immunogenicity. Bower et al. [2004] (adjuvant comparison, vaccine antigen design)

**No.** 963

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Subtype multiple

Neutralizing no

Immunogen vaccine

> Vector/Type: protein Strain: B clade IIIB, A clade UG37, B clade HAN2, D clade UG21, F clade BR29 HIV component: gp140, gp120ΔV1, V2, and V3 Adjuvant: Complete Freund's Adjuvant (CFA), Incomplete Freund's Adjuvant (IFA)

Species (Isotype) rabbit

References Jeffs et al. 2004

Keywords subtype comparisons, vaccine antigen design • A set of oligomeric envelope proteins were made from six primary isolates for potenial use as vaccine antigens: 92/UG/037 (clade A), HAN2/2 (clade B), 92/BR25/025 (clade C), 92/UG/021 (clade D), 93/BR/029 (clade F) and MVP5180 (clade O). This was one of a panel of MAbs used to explore folding and exposure of well characterized epitopes. The clade C isolate BR25 is apparently misfolded, as conformationdependent antibodies did not bind to it. Polyclonal sera raised in rabbits against the A, B, D and F antigens, which were deemed pure enough for immunization, as well as IIIB and IIIB with the V1, V2 and V3 loops deleted, cross-bound the other antigens, so shared epitopes across clades, but none of the sera had neutralizing activity. Jeffs et al. [2004] (vaccine antigen design, subtype comparisons)

**No.** 964

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Subtype B, CRF01\_AE

Neutralizing

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade MN, B clade GNE8, E clade CM244 HIV component: gp120 Adjuvant: aluminum hydrox-

Species (Isotype) human

References Lee et al. 2001

**Keywords** assay development, subtype comparisons, vaccine antigen design, vaccine-induced epitopes

• An assay was developed that characterizes antibody binding to primary isolates, and using this system there was a correlation between binding activity and neutralization by sera from HIV-infected people and gp120 vaccinated individuals. The magnitude and breadth of oligomeric, cell surface gp120 binding Abs induced by HIV-1 subtype B vaccines was characterized. The responses in people vaccinated with monoand bivalent rgp120 vaccines (AIDSVAX B and AIDSVAX B/B AIDSVAX B/E) indicated that increasing the number of antigens increased the cross-binding activities, in support of polyvalent vaccines. Lee et al. [2001] (assay development, vaccine antigen design, vaccine-induced epitopes, subtype comparisons)

No. 965 MAb ID polyclonal **HXB2 Location** Env **Author Location** Env **Epitope** Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: gp120-MAb A32 complex Strain: B clade 89.6, B clade BaL HIV component: gp120-Mab complex Adjuvant: Cholera toxin (CT), Complete Freund's Adjuvant (CFA), Incomplete Freund's Adjuvant (IFA), Ribi adjuvant (MPL+TDM) (RIBI)

Species (Isotype) guinea pig

References Liao et al. 2004

Keywords vaccine antigen design

• A32-rgp120 complexes opened up the CCR5 co-receptor binding site, but did not induce neutralizing antibodies with greater breadth among B subtype isolates than did uncomplexed rgp120 in vaccinated guinea pigs. The vaccine that gave the greatest breadth comparing A32-rgp120 BaL, A32-rgp120 89.6, rgp120 BaL, and rgp120 89.6, was the uncomplexed rgp120 BaL, as it neutralized 9/14 B clade isolates tested (60%). Liao et al. [2004] (vaccine antigen design)

No. 966

MAb ID polyclonal

HXB2 Location Env

Author Location gp120 (JRFL)

**Epitope** 

Subtype B

Neutralizing yes

Immunogen vaccine

Vector/Type: DNA Strain: B clade JRFL HIV component: gp120 Adjuvant: C3d fu-

sion

Species (Isotype) humanized mouse (IgG)

References Liu et al. 2004

Keywords adjuvant comparison

• BALB/c mice were immunized with codon-optimized or C3dfused DNA vaccine constructs and analyzed for their ability to elicit humoral and cell-mediated immune responses. Each strategy increased binding and gave rise to earlier appearance of neutralizing antibody respones against IIIB and MN viruses, but the combination did not act synergistically. C3d and codon optimization also gave enhanced CD8+ T cell responses to the epitope SIHIGPGRAFYTTGE. Liu et al. [2004] (adjuvant comparison)

No. 967

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env (SF2)

**Epitope** 

Subtype multiple

Neutralizing yes

Immunogen vaccine

Vector/Type: protein Strain: B clade SF2 HIV component: gp120 Adjuvant: aluminum hydroxide, Incomplete Freund's Adjuvant (IFA), MF59, Other

Species (Isotype) baboon

References Haigwood et al. 1992

**Keywords** adjuvant comparison, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-neutralization

• Baboons were given intramuscular immunization with env 2-3 SF2 (aa Ile-26 to Ala-510) or rgp120SF2. Native, glycosylated rgp120 SF2, gave a broader range of heterologous neutralizing Ab responses than denatured, non-glycosylated env 2-3 SF2. Repeated immunizations with the native rgp120 gave rise to weak but detectable NAbs against two African strains, NDK and ZR6. IFA/MTP-PE gave the highest titer antibodies.of many adjuvant combinations tested. Haigwood et al. [1992] (adjuvant comparison, vaccine antigen design, variant cross-recognition or cross-neutralization, vaccine-specific epitope characteristics)

No. 968

MAb ID polyclonal

**HXB2 Location** Env

Author Location Env (89.6)

Epitope Subtype B Neutralizing yes

Immunogen vaccine

Strain: B clade 89.6 HIV component: gp140, gp160,  $gp160\Delta V3$ ,  $gp140\Delta V3$ 

Species (Isotype) macaque, mouse References Lorin et al. 2004

**Keywords** vaccine antigen design, variant cross-recognition or cross-neutralization

• Mice susceptible to MV infection were intraperitoneally immunized with native HIV-1 89.6 env gp160 and gp140 and δV3 HIV-1 89.6 mutants expressed in live attenuated Schwarz measles vector (MV). The gp160ΔV3 construct raised more cross-reactive NAbs to primary isolates than did native gp160, and sera from the gp160ΔV3 animals neutralized SHIV 89.6, clade B strains Bx09, 92US660 and 92US714, and clade A virus 3253 but not to clade B 92HT593, at a 1:30 dilution. A HIVIG/2F5/2G12 combination was used as a positive control and could neutralize all isolates. The vaccine constructs had an additional 2F5 MAb epitope, ELDKWAS, but responses were not directed towards this epitope. Mice and macaques could raise anti-HIV responses in mice and macaques with preexisting MV immunity. Lorin et al. [2004] (vaccine antigen design, variant cross-recognition or cross-neutralization)

**No.** 969

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References McCaffrey et al. 2004

Keywords antibody binding site definition and exposure,

vaccine antigen design

• Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The loss of the glycan within the V3 loop (GM299 V3) and adjacnt to the C-terminal end of the V3 loop (GM329 C3) increased neutralization susceptibility to both sera, but the loss of sites in C2, C4, and V5 did not alter neutralization susceptibility. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey *et al.* [2004] (antibody binding site definition and exposure, vaccine antigen design)

**No.** 970

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp120 (HXBc2)

Epitope Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: Con A-NS Strain: B clade

HXBc2 HIV component: Env

Species (Isotype) macaque (IgA, IgG)

References Miyake et al. 2004

Keywords genital and mucosal immunity

Intranasal immunizations of three macaques with SHIV-nanospheres (SHIV-NS) induced vaginal anti-HIV-1 gp120 IgA and IgG antibodies. After intra-vaginal challenge with SHIV KU-2, 1/3 control animals and 1/3 SHIV vaccinated animals were infected, but the SHIV vaccinated animals had low viral loads that fell to undetectable levels. After intravenous re-challenge, all animals were infected, but SHIV immunized animals had lower viral loads. Miyake et al. [2004] (genital and mucosal immunity)

No. 971

MAb ID polyclonal

HXB2 Location Env

Author Location gp41 (HXB2)

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Opalka et al. 2004

**Keywords** assay development, assay standardization/improvement

An antigen panel representing different regions of gp41 was generated, and sera from 23 individuals were screened. Antigp41 titers were very high, and sera bound to many regions of gp41, there were no immunologically silent regions. Many individuals had broad responses to diverse regions. High titer responses tended to focus on the N-heptad, C-heptad and 2F5-4E10 regions, but there was no correlation between neutralization capacity of sera and the particular peptides recognized. Opalka et al. [2004] (assay development, assay standardization/improvement)

No. 972

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Research Contact Abraham Pinter, Public Health Research Institute, Newark, NJ, 07103. pinter@phri.o

References Pinter et al. 2004

Country United States

**Keywords** variant cross-recognition or cross-neutralization

• V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. 28 sera were tested - 24/28 sera gave greater than 90% neutralization of SF162 at dilutions of 1:180, while only 2/28 could give 90% neutralization of JRFL, and only 9/28 gave 50% neutralization at dilutions of 1:180. A chimera with SF162 V1V2 in a JRFL Env backbone was neutralization sensitive to most sera at a comparable level to SF162 Env, and in some cases the JRFL-SF162 V1V2 chimera was even more sensitive than JRFL. Pinter et al. [2004] (variant cross-recognition or cross-neutralization)

**No.** 973

MAb ID polyclonal

**HXB2 Location** Env

Author Location Env (gp160)

**Epitope** 

Subtype multiple

Neutralizing

Immunogen vaccine

Vector/Type: DNA, DNA prime with protein boost Strain: B clade LAI, A clade 92UG031, C clade 92BR025 HIV component on 160 Adjunction CM CSE

nent: gp160 Adjuvant: GM-CSF

**Species (Isotype)** mouse (IgG)

References Rollman et al. 2004

**Keywords** adjuvant comparison, enhancing activity, Th1, Th2, vaccine antigen design, variant cross-recognition or cross-neutralization

• Vaccination of mice with subtype B Env raised antibodies primarily against subtype B alone, while A+B+C clade Envs raised antibodies that could neutralize the autologous B, C strains, and weakly neutralize the A strain. Serum IgG responses to gp120s including all gp120 variable regions were induced in animals vaccinated with subtypes A, B and C of HIV-1 gp160 with rGM-CSF as adjuvant. Boosting with rgp160 with CpG-ODN enhanced IgG responses, shifted the Th1/Th2 to be more balanced, and these animals made both IgG and Ig2a responses and had expanded recognition of constant regions. The

B clade vaccine was LAI, and the A and C clade vaccines were actually V1-V5 of the A and C strains cloned into a LAI backbone. gp41 peptides were also recognized by sera. T cell responses to the multi-clade vaccine had enhanced cross-reactive CD4 T-cell proliferative responses, but diminished gamma IFN CD8 T-cell responses. Rollman *et al.* [2004] (adjuvant comparison, enhancing activity, vaccine antigen design, variant cross-recognition or cross-neutralization, Th1, Th2)

No. 974

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: DNA Strain: B clade IIIB HIV component: gp120 Adjuvant: C3d fu-

sion

Species (Isotype) mouse (IgG, IgG2a)

References Toapanta & Ross 2004

Keywords adjuvant comparison, Th1, Th2

• Mice [C57BL/6 (H-2b), BALB/c (H-2d), C3H/H3 (H-2k) and CD-1 Swiss] were vaccinated DNA carrying with 2 or 3 complement C3d genes fused to secreted sgp120. Responses were enhanced with C3d, particularly in outbred mice. sgp120-C3d-DNA vaccination induced a primarily IgG1 anti-Env Ab response in inbred mouse strains, while outbred mice had mixed IgG1/IgG2a responses; similarly IL4 (Th2) T-cell responses were observed in inbred mice, and mixed IL4 and IFN gamma (Th1/Th2) responses were observed in outbred mice. An increased avidity maturation of anti-Env Abs in outbred mice was also observed. Toapanta & Ross [2004] (adjuvant comparison, Th1, Th2)

No. 975

MAb ID polyclonal

HXB2 Location Env

**Author Location** gp120

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: canarypox prime with gp120 boost Strain: B clade LAI, B clade MN HIV component: Gag, gp120, Protease

Species (Isotype) human (IgA, IgG)

References Wright et al. 2004

**Keywords** genital and mucosal immunity, vaccine antigen design

HIV-1 specific responses were seldom detected after systemic
or mucosal vaccination with HIV gp120 in a canarypox vector
with a rgp120 boost. A limited IgA and CTL response was
observed after rectal vaccination, but overall, canary pox virus
was not an effective mucosal immunogen. Wright et al. [2004]
(genital and mucosal immunity, vaccine antigen design)

No. 976 MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env (735–752)

**Epitope Neutralizing** 

Immunogen vaccine

Vector/Type: peptide HIV component: gp41 Adjuvant: Complete Freund's Adjuvant

(CFA)

Species (Isotype) human, rabbit

References Kennedy et al. 1986

Keywords assay standardization/improvement

• Rabbits intramusculary immunized with peptide KLH ("HTLV-III aa 735-752") produced peptide-specific, serum Ab responses. In an ELISA, AIDS patient derived antisera tested positive for gp41-specific Ab. Kennedy et al. [1986] (assay standardization/improvement)

No. 977

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection, vaccine

Species (Isotype) human

References Zolla-Pazner 2004

Keywords review, vaccine antigen design

· This review summarizes neutralizing epitopes on Env and their use as vaccine antigens. Most antibodies are not neutralizing, and while some antibodies directed to conserved domains can neutralize the virus, these are generally poorly immunogenic. Variable loops do not elicit much cross-reactive neutralization, although the stem regions of these loops are more conserved so may have some promise. Polyclonal pooled sera from infected people can generally neutralize heterologous virus, suggesting that neutralizing epitopes are yet to be discovered. Polyvalent vaccine design is considered key. Zolla-Pazner [2004] (vaccine antigen design, review)

No. 978

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing yes

Immunogen vaccine

B clade IIIB HIV component: Strain: gp120, gp160 Adjuvant: aluminum hydrox-

ide

Species (Isotype) human, chimpanzee

References Berman et al. 1994

**Keywords** variant cross-recognition

neutralization

• Antisera derived from human or chimpanzee immunized with IIIB-rgp120 showed broad cross-reactivity to HIV-1 isolates MN, IIIB, JRcsf and NY-5 (subtype B), Z6 (subtype D), A244 (subtype E)and Z321 (subtype A). Sera of IIIB-rgp120 chimpanzees cross-reacted with 6/8 V3 peptides derived from HIV-1 isolates (MN, NY5, SF2, RF, CDC4 and IIIB). Human sera

only recognized 1/8 V3 peptides, HIV-1 MN. The magnitude, duration, avidity and half-life of IIIB-rgp120-specific Ab-responses were species specific. Sera derived from IIIBrgp120-immunized humans and chimpanzees inhibited binding of both IIIB- and MN-derived rgp120 to cell-surface CD4. Berman et al. [1994] (variant cross-recognition or crossneutralization)

No. 979

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (V3)

**Epitope** 

Subtype A

**Neutralizing** 

Immunogen vaccine

Vector/Type: hepatitis B surface antigen lipoprotein particles (HsBAg) Strain: A clade HIV component: CD4BS, V3 Adjuvant: Complete Freund's Adjuvant (CFA), Incomplete Freund's Adjuvant (IFA)

Species (Isotype) mouse

References Cruz et al. 2004

Keywords variant cross-recognition crossneutralization

• Vaccinations with either the subtype A V3 consensus in a Hepatitis B carrier protein, or a multiple antigen peptide (MAP) construct (mixotope) carrying some 5000 V3s representing the diversity of subtype A V3 loops, were compared. Each was combined with a C4 peptide spanning a region involved in CD4 binding. BALB/c mice were used for immunization. The consensus V3 gave higher and more cross-reactive responses than the mixotope. The mixotope response was restricted to the most conserved region of V3, while the consensus antibodies tended to recognize heptamers representing both sides and the tip of the loop. Antibodies against the C4 region were also raised. Cruz et al. [2004] (variant cross-recognition or cross-neutralization)

No. 980

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: DNA with CMV promotor, modified vaccinia Ankara (MVA) Strain: B clade IIIB HIV component: gp120 Adjuvant: Cholera toxin (CT)

Species (Isotype) mouse (IgA, IgG, IgG1)

References Gherardi et al. 2004

Keywords adjuvant comparison, genital and mucosal immunity

Env-specific IgG and IgA Abs were detected in vaginal washings of BALB/c mice (H-2Dd) following intranasal immunization of rMVA + CT in both MVA/MVA and DNA/MVA schemes. Coadministration of CT as adjuvant resulted in an

increased responses. Gherardi *et al.* [2004] (**adjuvant comparison**, **genital and mucosal immunity**)

**No.** 981

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype B

Neutralizing yes

Immunogen SHIV infection

Species (Isotype) macaque

References Montefiori et al. 1998

**Keywords** variant cross-recognition or crossneutralization

 Neutralizing antibody responses in rhesus macaques infected with SHIV variants HXB2, 89.6, and 89.6PD were studied. The SHIV infections resulted in induction of high-titer neutralizing Abs to homologous SHIV and HIV-1 strains; heterologous NAbs responses were infrequent and only detected after 40 weeks of infection. Montefiori *et al.* [1998] (variant crossrecognition or cross-neutralization)

No. 982

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp160 (IIIB)

Epitope Subtype B Neutralizing yes Immunogen vaccine

Vector/Type: DNA prime with gp160 boost Strain: B clade IIIB HIV component: gp160 Adjuvant: Incomplete Freund's Adjuvant

(IFA), IL-12

Species (Isotype) macaque

References Rasmussen et al. 2002

**Keywords** adjuvant comparison, vaccine-induced epitopes

· DNA prime vaccinations by intradermal or gene-gun delivery were given to neonatal macaques, with or without IL-12, followed by boosting with gp160, or else gp160 was given without the DNA prime. Many of the animals had neutralizing antibodies against autologous Env, and no CTL was detected prior to challenge after DNA inoculation. Autologous SHIV-vpu+ challenge was contained in 4/15 DNA prime-gp160 boost-vaccinated macaques and in 3/4 animals only receiving gp160. Six animals that contained virus were rechallenged with autologous virus, and the virus was rapidly cleared. After an additional challenge with heterologous pathogenic SHIV 89.6P, 4/6 maintained low or limited viral infection and normal CD4 counts. Two animals gave evidence of Gag specific CTL and proliferative responses after pathogenic SHIV challenge with 89.6P, with no other evidence of infection. Rasmussen et al. [2002] (adjuvant comparison, vaccine-induced epitopes)

No. 983 MAb ID polyclonal HXB2 Location Env

Author Location gp140 (SF162)

Epitope Subtype B Neutralizing yes

Immunogen vaccine

*Vector/Type:* DNA prime with protein boost *Strain:* B clade SF162 *HIV component:* 

gp140, gp140 $\Delta$ V2

Species (Isotype) macaque

**Ab Type** gp120 C5, gp120 C1-C2, gp120 CD4BS,

gp120 V3, gp120 V1-V2

References Srivastava et al. 2003

Keywords vaccine antigen design, vaccine-induced epi-

topes

• Vaccination of macaques with SF162gp140 was compared with vaccination with SF162 deltaV2gp140. V1, V2, V3, CD4BS, C1 and C2 antibodies were elicited by the intact SF612, and the antibodies were able to neutralize some heterologous strains. Deletion of the V2 loop altered the response so that there was a higher ratio of CD4BS antibody made relative to V3 antibody, but did not increase overall amount of CD4BS antibodies. Antibodies against C5 were also elicited by the deltaV2 construct. Overall, the deltaV2 construct was better able to raise antibodies that could cross-neutralize heterologous strains. Using a cleaved versus fused form of gp120 altered the ratio of C1 to C5 antibodies raised, with more C5 response to the fused form. Srivastava et al. [2003] (vaccine antigen design, vaccine-induced epitopes)

No. 984

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120 (SF2)

Epitope Subtype B Neutralizing yes

Immunogen HIV-1 infection, vaccine

Vector/Type: protein Strain: B clade SF2 HIV component: gp120 Adjuvant: aluminum hydroxide, Incomplete Freund's Adjuvant (IFA), muramyl-dipeptide base adjuvant

(Syntex)

Species (Isotype) human, baboon

References Steimer & Haigwood 1991

Keywords adjuvant comparison, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-neutralization

• Immunization with native glycosylated rgp120SF2 produced Abs directed against linear and conformational epitopes. Denatured, deglycosylated env2-3 (SF2) produced Abs against only linear determinants. Sera from 8/8 rgp120SF2 vs 3/8 env2-3SF2 immunized baboons cross-neutralized HIV-MN. Only 5/8 rgp120SF2 vaccinated animals had neutralizing activity against HIV-HTLV-IIIB and HIV-BRU. Abs from infected people who reacted with rgp120SF2 showed broad cross-neutralization of HIV-1MN, HIV- BRU, HIV-Zr6 and HIV-SF2 isolates, in comparison to env2-3(SF2)immunization, which only neutralized HIV-1 MN. Stronger neutralization potency of Ab responses was observed in baboons using Alum and MF101 as adjuvants. Steimer & Haigwood [1991] (adjuvant comparison,

vaccine antigen design, variant cross-recognition or cross- Species (Isotype) macaque (IgG) neutralization, vaccine-specific epitope characteristics)

No. 985

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype)

References Balzarini 2005

Keywords antibody binding site definition and exposure · Author hypothesizes that resistance to drugs that target glycosylation sites of gp120 might stimulate deletions within the glycan shield, thus exposing novel epitopes that enhance neutralization susceptibility. Balzarini [2005] (antibody binding site definition and exposure)

No. 986

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Subtype A

**Neutralizing** yes

Immunogen vaccine

*Vector/Type:* virus-like particle (VLP)

Strain: A clade 94UG018 HIV component:

anchored gp120

Species (Isotype) mouse (IgA, IgG)

References Buonaguro et al. 2005

Keywords vaccine antigen design, variant crossrecognition or cross-neutralization

• The impact of vaccination routes was studied in BALB/c mice for clade A gp120 in VLPs. I.n. and i.p. vaccination gave a systemic and mucosal IgG and IgA response, and a CTL response. Higher specific IgA titers were detected in i.n. vaccinated mice, and CTL responses were stronger in the i.p. group. The oral route did not induce NAb responses. gp120-Env (V3 loop, TR-PYNNTRQSTHIGPGQALYTTNIIGDIRQAHC)specific IgG and IgA Ab were detected at 1-2-dilution lower dilutions than p24 Abs. Neutralizing activity (>50%) against the autologous clade A Ugandan and a heterologous clade B Italian field isolate was observed. No adjuvants were used. Buonaguro et al. [2005] (vaccine antigen design, variant cross-recognition or cross-neutralization)

No. 987

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Subtype CRF02\_AG

Neutralizing no

Immunogen vaccine

Vector/Type: virus-like particle (VLP), DNA prime with modified vaccinia Ankara (MVA) boost Strain: CRF02 IC0928 HIV compo-

nent: Env, Gag, Pol

References Ellenberger et al. 2005

Keywords vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition

or cross-neutralization

 Macaques were given a Gag-Pol-Env DNA prime followed by a MVA boost, comparing two DNA constructs, one that resulted mature VLPs with processed Gag (IC48) and one that had a point mutation in Gag that resulted in immature VLPs (IC1-90). Both vaccines gave antibody and T-cell responses to Env and Gag, although negligible neutralizing antibody responses were found. Autologous virus is difficult to neutralize, but the antibodies in the vaccinated macaques also did not neutralize the laboratory adapted B clade MN strain. Ellenberger et al. [2005] (vaccine antigen design, variant cross-recognition or crossneutralization, vaccine-specific epitope characteristics)

No. 988

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Subtype A, B, C

Neutralizing P

Immunogen vaccine

Vector/Type: DNA, adenovirus Strain: B clade HXB2, A clade 92RW020, C clade 97ZA012 HIV component: gp140ΔCFI

Species (Isotype) guinea pig

Ab Type gp120 V3

References Chakrabarti et al. 2005

Keywords vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition

or cross-neutralization

· Guinea pigs were immunized with a hybrid HXB2/BaL Env (HIV HXB/BaL gp140 $\delta$ CFI, clade B) in which the tip of the V3 loop (GPGRA) was replaced with the 2F5 epitope LELD-KWAS. 2F5 bound to the Env that carried the V3-replacement 2F5 epitope, but antibodies against this construct only neutralized the X4-tropic lab adapted HIV strain IIIB, and not CCR5-HIV BaL or SF162 isolates. This immunogen, a single B clade immunogen, and a mixture of A + B + C clade envelopes, were compared. The single B clade immunogen had neutralizing activity against some B clade viruses. The A+B+C mixture was found to maintain the B clade responses, while eliciting NAbs with greater breadth when tested against a panel 19 A, B and C clade primary isolates. Chakrabarti et al. [2005] (vaccine antigen design, variant cross-recognition or crossneutralization, vaccine-specific epitope characteristics)

No. 989

MAb ID polyclonal

**HXB2 Location** Env Author Location gp120

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: virus-like particle (VLP) Strain: B clade BaL HIV component: Env, Gag Adjuvant: block copolymer CRL8623

Species (Isotype) guinea pig

References Hammonds et al. 2005

**Keywords** adjuvant comparison, vaccine antigen design

· Adjuvanted (either with a block copolymer or with a CpG aluminum hydroxide adjuvant) pseudovirions and with a recombinant gp120 boost gave significant gp120-specific NAb responses to autologous virus relative to pesudoviruses alone and to 2/5 additional primary isolates (SS1196 and Pvo) tested. Hammonds et al. [2005] (adjuvant comparison, vaccine antigen design)

**No.** 990

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: gp120 Adjuvant: C3d fusion, Ribi adjuvant

(MPL+TDM) (RIBI)

Species (Isotype) mouse (IgG)

References Koch et al. 2005

Keywords adjuvant comparison

• Fusion of C3d repeats and the addition of Ribi adjuvant to gp120 variant glycoprotein (gp120δC1/C5(C3d)2,enhanced gp120-specific Ab responses, but Ribi alone gave almost comparable enhancement. Thus C3d as an adjuvant may be of particular value when used alone in conditions where avoiding denaturation and preservation of the native structure is important. Koch et al. [2005] (adjuvant comparison)

No. 991

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** Env (JRFT)

**Epitope** Subtype B **Neutralizing** 

Immunogen vaccine

Vector/Type: adenovirus Strain: B clade

JRFL HIV component: Gag, gp140

Species (Isotype) macaque

References Liang et al. 2005

Keywords vaccine antigen design, vaccine-induced epitopes

• 4/4 Mamu-A\*01-negative rhesus monkeys that were vaccinated with gp140 and challenged intravenously with SHIV-89.6P produced significant neutralizing Ab titers by day 28 realtive to other challenge groups, even though no pre-challenge NAb was detected, suggesting the existance of prechallenge memory neutralizing Ab responses. The viral set point was associated with the strength of the cellular immune response to Gag and Env, but not to Tat. Liang et al. [2005] (vaccine antigen design, vaccine-induced epitopes)

No. 992

MAb ID polyclonal

**HXB2 Location** Env

**Author Location (MN)** 

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Raviv et al. 2005

Keywords vaccine antigen design

Retrovirus inactivation for vaccine antigen delivery was explored through lipid modification by hydrophobic photoinduced alkylating probe 1.5 iodonaphthylazide (INA). The viral proteins were shown to be structurally intact in the treated noninfectious virus, through the preservation of antibody binding sites for polyclonal anti-gp120 serum, and for broadly neutralizing MAbs 2G12, b12 and 4E10, although the modifications of the lipid disabled viral infection. Raviv et al. [2005] (vaccine antigen design)

No. 993

MAb ID polyclonal

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen

Species (Isotype)

References Rossi et al. 1989

Keywords mother-to-infant transmission

• Abs that bound to gp120 peptides were found to correlate with lack of transmission in infants less than 6 months old and born to HIV+ mothers. Maternal Abs to these same peptides were also enriched in mothers that did not transmit. Rossi et al. [1989] (mother-to-infant transmission)

No. 994

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide Adjuvant: gp41 N-HR

and C-HR helical peptides

Species (Isotype) rabbit (IgG)

**Ab Type** C-HR, N-HR, gp41six-helix bundle

References Golding et al. 2002b; de Rosny et al. 2001

- The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry - preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-sixhelix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion – the preincubation 31.5 C step did not alter anti-C-HR Abs inability to inhibit fusion. Golding et al. [2002b]
- A panel of Abs against gp41 heptad repeats N-HR, C-HR, and self-assembled stable N-HR and C-HR six helix bundles were generated. de Rosny et al. [2001]

No. 995

MAb ID 101-342

**HXB2 Location** Env

Author Location gp120 (476–505 HAM112, O group)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: O group HAM112 HIV component: gp160

**Species (Isotype)** mouse (IgG2a $\kappa$ )

Ab Type C-term

References Scheffel et al. 1999

• 101-342: Overlapping peptides based on group O HAM112 Env were tested for MAb reactivity. Scheffel *et al.* [1999]

No. 996

**MAb ID** 101-451

**HXB2 Location** Env

Author Location gp120 (498–527 HAM112, O group)

Epitope Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: O group HAM112 HIV component: gp160

**Species** (**Isotype**) mouse ( $IgG2b\kappa$ )

Ab Type C-term

References Scheffel et al. 1999

• 101-451: Overlapping peptides based on group O HAM112 Env were tested for MAb reactivity. Scheffel *et al.* [1999]

No. 997

MAb ID 120-1

**HXB2 Location** Env

**Author Location** gp120 (503-532)

**Epitope Neutralizing** no

Immunogen vaccine

Vector/Type: peptide

**Species** (**Isotype**) mouse ( $IgM\kappa$ )

Ab Type C-term

References Dalgleish et al. 1988; Chanh et al. 1986

No. 998

MAb ID T26

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein

Species (Isotype) mouse

Ab Type C-term

Research Contact Patricia Earl, National Institute of Allergy and

Infectious Diseases

References Kilgore et al. 2003; Earl et al. 1997; Earl et al.

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Keywords antibody binding site definition and exposure,

antibody generation, variant cross-recognition

or cross-neutralization

• T26: Mab is restricted in its binding to gp41 of the LAI isolate and not to gp41 of the MN, Ada and RF isolates. Antibody specificity may be determined by LAI residues D637E, N641D and H648Y. T26 binds to the N-terminal half of the C helix (aa630-680) of the LAI envelope, specifically targeting a confomational epitope within the six-helix bundle of gp41. Addition of the C-helical peptide inhibitor from LAI (T26 reactive) rescued the binding activity of MAb T26 to cell-surface expressed RF envelope (T26 non-reactive) triggered with sCD4 or cell-surface expressed receptors in a surface immunoprecipitation assay. This supports that C-peptide entry inhibitors bind to the gp41 N-helical coiled-coil, disrupting native six-helix bundles. Kilgore et al. [2003] (antibody binding site definition and exposure)

- T26: T26 was raised against the gp140 tetramer, binds to gp41 and is a highly strain specific. Earl *et al.* [1997] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- T26: A panel of 138 MAb raised against different forms of soluble Env. Earl *et al.* [1994] (antibody generation)

No. 999

MAb ID D33

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS, C-term, N-term

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- D33: A comparison of 25 gp120 specific, conformation dependent MAbs was done D33 is part of a group of MAbs labeled A1 all A1 MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, blocked CD4 binding, were sensitive to mutations in gp120 positions 368 and 370 that directly contact CD4 D33 was unusual for the group of A1 MAbs, because while it blocked CD4 binding completely, but competed with MAbs that did not in a BIAcore assay both the N- and C-terminal ends of gp120 are involved in D33 binding. Sugiura et al. [1999]
- D33: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

**No.** 1000

MAb ID polyclonal

HXB2 Location Env

**Author Location** 

Epitope

Neutralizing

Immunogen HIV-1 infection Species (Isotype) human (IgA)

Ab Type gp120 CD4BS, C-term, gp120 V3-C4

References Vincent et al. 2004

Country France

## **Keywords** genital and mucosal immunity

• IgA derived from sera and saliva from 5 HIV-1 infected patients undergoing ART therapy reacted to peptide antigens corresponding to the C3-V4 region of gp120 and the C-terminal part of gp41. HIV-1-specific IgA obtained in 6/26 sera and 5/25 saliva samples inhibited gp120-sCD4 protein binding. Vincent et al. [2004] (genital and mucosal immunity)

**No.** 1001 MAb ID 212A **HXB2 Location** Env Author Location gp120 **Epitope** Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 C1

Research Contact James Robinson, Tulane University, LA

References Pantophlet et al. 2004; Pantophlet et al. Research Contact G. Robey, Abbott Inc. 2003b; Binley et al. 1998; Sullivan et al. 1998b; Parren et al. 1997b; Wyatt et al. 1997; Ditzel et al. 1997; Fouts et al. 1997; Binley et al. 1997a; Moore & Sodroski 1996; Moore et al. 1994d; Robinson et al. 1992

## Keywords vaccine antigen design

- 212A: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including 212A. Pantophlet et al. [2004] (vaccine antigen design)
- 212A: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional Nlinked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)
- 212A: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer. Binley et al.
- 212A: Does not compete with binding of MAb generated in response to gp120-CD4 complex, CG10. Sullivan et al. [1998b]
- 212A: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding – 212A bound monomer, did not bind oligomer or neutralize JRFL. Fouts et al. [1997]
- 212A: Does not neutralize TCLA strains or primary isolates. Parren et al. [1997b]
- 212A: Binds efficiently to sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding – does

not bind to HXBc2 gp120 if the 19 C-term amino acids are deleted. Wyatt et al. [1997]

- 212A: Binding enhanced by anti-V3 MAb 5G11 reciprocal inhibition with anti-C1 MAbs. Moore & Sodroski [1996]
- 212A: Mutations that inhibit binding: C1 (45 W/S) and V5 (463 N/D) - and enhance binding: V2 (179/180 LD/DL) and C5 (495 G/K). Moore et al. [1994d]

No. 1002 MAb ID 522-149 **HXB2 Location** Env Author Location gp120 **Epitope** 

Neutralizing no Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) mouse Ab Type gp120 C1

References Pantophlet et al. 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Yang et al. 2000; Binley et al. 1998; Trkola et al. 1996a; Moore & Sodroski 1996

Keywords antibody interactions, vaccine antigen design

- 522-149: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including 522-149. Pantophlet et al. [2004] (vaccine antigen design)
- 522-149: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)
- 522-149: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a C1-binding Fab that had no impact on 4KG5 binding. Zwick et al. [2003] (antibody interactions)
- 522-149: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes – CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 - non-neutralizing MAbs C11, A32,

522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 – MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang *et al.* [2000]

- 522-149: A panel of MAbs were shown to bind with similar
  or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta
  V1, V2, and V3), thus such a core protein produces a structure
  closely approximating full length folded monomer. Binley et al.
  [1998]
- 522-149: Binding is enhanced by C5 antibodies M91 and 1C1

   mutual binding-inhibition with anti-C1 antibody 133/290 –
   binding is destroyed by a W/L (position 61, LAI) gp120 amino acid substitution other C1 antibodies enhance binding to gp120. Moore & Sodroski [1996]
- 522-149: Does not neutralize JR-FL nor block gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]

**No.** 1003

MAb ID CA1 (ARP3117)

HXB2 Location Env Author Location Env Epitope Subtype A Neutralizing

Immunogen vaccine

Vector/Type: vaccinia prime with gp120 boost Strain: A clade HIV component: Env

Species (Isotype) mouse Ab Type gp120 C1 References Jeffs et al. 2004

**Keywords** subtype comparisons, vaccine antigen design • CA1: A set of oligomeric envelope proteins were made from six primary isolates for potenial use as vaccine antigens: 92/UG/037 (clade A), HAN2/2 (clade B), 92/BR25/025 (clade C), 92/UG/021 (clade D), 93/BR/029 (clade F) and MVP5180 (clade O). This was one of a panel of MAbs used to explore folding and exposure of well characterized epitopes. The clade C isolate BR25 is apparently misfolded, as conformationdependent antibodies did not bind to it. CA1 is a MAb that binds to a linear epitope in the C1 region of gp120 that was raised against clade A variant 92/UG/029. CA1 was subtypespecific and bound only to the antigen from all clade A. Polyclonal sera raised in rabbits against these antigens cross-bound the other antigens, but none of the sera had neutralizing activity. Jeffs et al. [2004] (vaccine antigen design, subtype comparisons)

**No.** 1004

**MAb ID** CA13 (ARP3119)

HXB2 Location Env Author Location Env Epitope Subtype A Neutralizing

Immunogen vaccine

Vector/Type: vaccinia prime with gp120 boost Strain: A clade HIV component: Env

Species (Isotype) mouse

Ab Type gp120 C1

References Zipeto et al. 2005; Jeffs et al. 2004

Keywords subtype comparisons, vaccine antigen design

- CA13: MRC Centralized Facility for AIDS Reagents, NIBSC, UK, ARP3119.
- CA13: HIV-1 fusion complexes were prepared from cell lines expressing R5 HIV-1 gp120/gp41 and CD4-CCR5. Neutralizing Abs were raised against both R5 (strain BaL) and X4 (strain 213) viruses. CA13 was used to detect gp120/gp41. Zipeto et al. [2005] (vaccine antigen design)
- CA13: A set of oligomeric envelope proteins were made from six primary isolates for potential use as vaccine antigens: 92/UG/037 (clade A), HAN2/2 (clade B), 92/BR25/025 (clade C), 92/UG/021 (clade D), 93/BR/029 (clade F) and MVP5180 (clade O). This was one of a panel of MAbs used to explore folding and exposure of well characterized epitopes. The clade C isolate BR25 is apparently misfolded, as conformation-dependent antibodies did not bind to it. CA13 is a MAb that binds to a linear epitope in the C13 region of gp120 that was raised against clade A variant 92/UG/029. C13 bound to antigens from all clades A-F, as well as group O. Polyclonal sera raised in rabbits against these antigens cross-bound the other antigens, but none of the sera had neutralizing activity. Jeffs et al. [2004] (vaccine antigen design, subtype comparisons)

**No.** 1005

MAb ID L19

**HXB2 Location** Env

Author Location gp120 (HXBc2)

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 C1

**References** Ditzel *et al.* 1997

• L19: gp120 immobilized on solid phase by capture with anti-CD4 BS MAb L72 was used for the selection of Fabs – six N-term Fabs, L19 L34, L35, L52, L59, and L69, were obtained that have a similar epitope to Fab p7. Ditzel *et al.* [1997]

**No.** 1006

MAb ID M90

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein HIV component: Env

Species (Isotype) (IgG1)

Ab Type gp120 C1

Research Contact Fulvia di Marzo Veronese

References Koefoed et al. 2005; Pantophlet et al. 2003b;

Yang *et al.* 2000; Binley *et al.* 1999; Binley *et al.* 1998; Wyatt *et al.* 1997; Ditzel *et al.* 1997; Moore & Sodroski 1996; DeVico *et al.* 

1995; di Marzo Veronese et al. 1992

Keywords antibody binding site definition and exposure

- M90: IgG antibody phage display libraries were created from HIV-1 + individuals after pre-selection of PBMC with gp120, as an alternative to using bone marrow for generating libraries.
   M90 was among a set of Abs used for competition studies to define the binding sites of the newly isolated MAbs, and has a conformational C1 epitope. Koefoed *et al.* [2005] (antibody binding site definition and exposure)
- M90: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of non-neutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet *et al.* [2003b]
- M90: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang *et al.* [2000]
- M90: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits – a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen - SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 - nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 - MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999]
- M90: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer. Binley *et al.* [1998]
- M90: Binds efficiently to sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding does not bind to HXBc2 gp120 if the 19 C-term amino acids, in conjunction with C1 positions 31-82, are deleted. Wyatt *et al.* [1997]
- M90: Reciprocal inhibition of binding of other anti-C1 MAbs

   inhibits CD4 binding site MAbs enhances binding of V2
   MAbs G3-4 and SC258. Moore & Sodroski [1996]

M90: Reacted with both non-reduced (but not denatured) covalently cross-linked gp120-CD4 complex. DeVico et al. [1995]

M90: Reactive only with native gp120, so binds to a discontinuous epitope – reacts with multiple strains. di Marzo Veronese et al. [1992]

**No.** 1007

MAb ID MAG 104

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain:

B clade HXB2  $\,\,$  HIV component: gp120  $\,$ 

Species (Isotype) mouse

Ab Type gp120 C1

Research Contact C. Y. Kang, IDEC Inc

References Kang et al. 1994

 MAG 104: Only observed amino acid substitution that reduces binding: 88 N/P and 106 E/A – does not bind to C1 region 20 mer peptides, tentative classification conformationally sensitive anti-C1 MAb. Kang et al. [1994]

**No.** 1008

MAb ID MAG 45 (#45, MAG45)

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

*Vector/Type:* sCD4-gp120 complex *Strain:* 

B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 C1

Research Contact C. Y. Kang, IDEC Inc, or Dr. Hariharam,

IDEC Pharmaceuticals Corporation, La Jolla,

CA

References Koefoed et al. 2005; Yang et al. 2000; Wyatt

et al. 1997; Moore & Sodroski 1996; Kang

et al. 1994

**Keywords** antibody binding site definition and exposure

- MAG 45: IgG antibody phage display libraries were created from HIV-1 + individuals after pre-selection of PBMC with gp120, as an alternative to using bone marrow for generating libraries. MAG 45 was among a set of Abs used for competition studies to define the binding sites of the newly isolated MAbs, representing a MAb with a C1 epitope. Koefoed *et al.* [2005] (antibody binding site definition and exposure)
- MAG 45: Called #45 a combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 MAbs directed at the extreme termini of gp120 C1 (135/9)

and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang et al. [2000]

- MAG 45: Called #45 binds to efficiently sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding – does not bind to HXBc2 gp120 if the 19 C-term amino acids, in conjunction with C1 positions 31-50, are deleted. Wyatt et al. [1997]
- MAG 45: Reciprocal binding inhibition with anti-C1-C5 and anti-C1-C4 discontinuous MAbs - binding enhanced by anti-V3 5G11 – inhibits binding of anti-CD4 binding site MAbs. Moore & Sodroski [1996]
- MAG 45: Only observed amino acid substitution that reduces binding: 88 N/P – does not bind to C1 region 20 mer peptides, tentative classification conformationally sensitive anti-C1 MAb. Kang et al. [1994]

No. 1009

MAb ID MAG 95

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain:

B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 C1

Research Contact C. Y. Kang, IDEC Inc

References Kang et al. 1994

• MAG 95: Only observed amino acid substitution that reduces binding: 88 N/P – does not bind to C1 region 20 mer peptides, tentative classification conformationally sensitive anti-C1 MAb. Kang et al. [1994]

**No.** 1010

MAb ID MAG 97

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

*Vector/Type:* sCD4-gp120 complex *Strain:* B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 C1

Research Contact C. Y. Kang, IDEC Inc

References Kang et al. 1994

• MAG 97: Only observed amino acid substitution that reduces binding: 88 N/P - does not bind to C1 region 20 mer peptides, Research Contact Patricia Earl and Christopher Broder, NIH tentative classification conformationally sensitive anti-C1 MAb. Kang et al. [1994]

**No.** 1011

MAb ID P35

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype) human

Ab Type gp120 C1

**References** Zwick et al. 2003: Kwong et al. 2002

Keywords antibody binding site definition and exposure, antibody interactions

- P35: called p35. scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restrict CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a C1-binding Fab with a discontinuous epitope that had no impact on 4KG5 binding. Zwick et al. [2003] (antibody interactions)
- P35:Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-299) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Authors describe the epitope as N-terminal, linear. Kwong et al. [2002] (antibody binding site definition and exposure)

No. 1012

MAb ID T9

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

Neutralizing

Immunogen vaccine

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 C1

References Golding et al. 2002b; Earl et al. 1997; Broder

et al. 1994

**Keywords** antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization

- There are two HIV-Abs with the name T9, one binds to gp41, one to gp120.
- T9: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry – preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and

anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion – the preincubation 31.5 C step did not alter the inhibitory activity of neutralizing Abs anti-gp41 2F5, or anti-gp120 2G12, IG1b12, 48d, and 17b - nor did it alter two gp41 MAbs, T9 and D61, inability to inhibit fusion. Golding et al. [2002b] (antibody binding site definition and exposure)

- T9: This antibody, along with 7 others (M10, D41, D54, T6, T4, T10 and T35), can block the linear murine MAb D61, and the human MAb 246-D, which both bind to the immunodominant region near the two Cys in gp41 – most of these antibodies are oligomer dependent – all of the MAbs are reactive with ten different HIV-1 strains – members of this competition group are blocked by sera from HIV-1 + individuals. Earl et al. [1997] (antibody binding site definition and exposure)
- T9: One of five MAbs (T4, T6, T9, T10 and T35) in a competition group that bind to a conformation-dependent epitope in gp41 and is oligomer specific - neutralizes IIIB and SF2. Broder et al. [1994] (antibody generation, variant crossrecognition or cross-neutralization)

**No.** 1013

MAb ID p7

**HXB2 Location** Env

Author Location gp120 (HXBc2)

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 C1

References Parren et al. 1997b; Ditzel et al. 1997

- p7: gp120 immobilized on solid phase by capture with sCD4 was used for selection of Fabs - three novel N-term Fabs were obtained that bind to similar epitopes, p7, p20, and p35 - a C1 W/S substitution at position 45 abolished binding, a Y/D at position 45 reduced binding, and C5 region substitutions 475 M/S and 493 P/K enhanced binding – compete with MAbs M85, M90 and 212A, but not M91 and G3-299. Ditzel et al.
- p7: Does not neutralize TCLA strains or primary isolates. Par- Species (Isotype) human ren et al. [1997b]

**No.** 1014

MAb ID L100

**HXB2 Location** Env

Author Location gp120 (HXBc2)

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

**Ab Type** gp120 C1-C2

References Kwong et al. 2002; Parren & Burton 1997; Parren et al. 1997b; Ditzel et al. 1997

Keywords antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization

• L100: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb

ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-299) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Authors describe the epitope as N-terminal, discontinuous. Kwong et al. [2002] (antibody binding site definition and exposure)

- L100: gp120 immobilized on solid phase by capture with sCD4 and then masked with Fab p7 allowed selection of a new Fab, L100, with a novel specificity for C1 and C2 - gp120 C1 substitutions 69 W/L and 76 P/Y abolish L100 binding, and C2 substitutions 252 R/W, 256 S/Y, 262 N/T and 267 E/L abolish or strongly inhibit L100 binding - inhibits binding of MAbs M90 and G3-299, but not M85, 212A, and M91. Ditzel et al. [1997]; Parren & Burton [1997] (antibody binding site definition and exposure, antibody generation)
- L100: Does not neutralize TCLA strains or primary isolates. Parren et al. [1997b] (variant cross-recognition or crossneutralization)

No. 1015

**MAb ID** 2/11c (211c, 2.11c, 211/c, 2-11c)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L (weak)

Immunogen HIV-1 infection

**Ab Type** gp120 C1-C4

Research Contact James Robinson, Tulane University, LA

References Kwong et al. 2002; Xiang et al. 2002a; Binley

et al. 1998; Wyatt et al. 1997; Li et al. 1997; Fouts et al. 1997; Binley et al. 1997a; Trkola

et al. 1996a; Moore & Sodroski 1996

**Keywords** antibody binding site definition and exposure • 2/11c: Called 211/c. Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-299)

MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Authors describe the epitope as N-terminus, discontinuous. Kwong et al. [2002] (antibody binding site definition and exposure)

- 2/11c: Used as a negative control in a study of CD4i MAbs. Xiang *et al.* [2002a]
- 2/11c: Called 211/c a panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein (Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer. Binley *et al.* [1998]
- 2/11c: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric env binding - 2/11c bound monomer, did not bind oligomer or neutralize JRFL. Fouts et al. [1997]
- 2/11c: Called 2.11c One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env – 50% neutralization could not be achieved at a maximal concentration of 67 mug/ml. Li et al. [1997]
- 2/11c: Binds efficiently to sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding – does not bind to HXBc2 gp120 if the 19 C-term amino acids, in conjunction with C1 positions 31-74, are deleted. Wyatt et al. [1997]
- 2/11c: Inhibits binding of anti-C1, -C5, -C4, -V3 and anti-CD4 binding site MAbs – induces binding of some anti-V2 and CD4i MAbs (48d and 17b) – similar reactivity pattern to A32, but less cross-reactive and lower affinity – A32 and 211/c are unique among known human and rodent MAbs. Moore & Sodroski [1996]
- 2/11c: Called 211c does not neutralize JR-FL nor block gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]

**No.** 1016

MAb ID A32

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1)

**Ab Type** gp120 C1-C4, gp120 adjacent to CD4BS

Research Contact James Robinson, Tulane University, New Or-

leans, LA, USA

References Selvarajah et al. 2005; Haynes et al. 2005; Pantophlet et al. 2004; Liao et al. 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Kwong et al. 2002; Grundner et al. 2002; Yang et al. 2002; Finnegan et al. 2001; Yang et al. 2000; Binley et al. 1999; Binley et al. 1998; Sullivan et al. 1998b; Parren et al.

1997b; Boots et al. 1997; Wyatt et al. 1997; Burton & Montefiori 1997; Fouts et al. 1997: Binley et al. 1997a; Trkola et al. 1996a; Wu et al. 1996; Moore & Sodroski 1996; Moore & Ho 1995; Wyatt et al. 1995; Moore et al. 1994b

Keywords antibody binding site definition and exposure, antibody interactions, co-receptor, mimotopes, review, subtype comparisons, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-neutralization

- A32: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)
- A32: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. V1/V2/V3 MAb 4KG2, C1-C4 MAb A32, C1-C5 MAb C11, and HIVIG all either did not bind or had significantly diminished binding to both antigen constructs. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)
- A32: A32-rgp120 complexes opened up the CCR5 co-receptor binding site, but did not induce neutralizing antibodies with greater breadth among B subtype isolates than did uncomplexed rgp120 in vaccinated guinea pigs. Liao et al. [2004] (vaccine antigen design)
- A32: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including A32. Pantophlet et al. [2004] (vaccine antigen design)
- A32: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional Nlinked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)
- A32: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results

suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. A32 is described as having a C1-C4 discontinuous CD4i epitope, and had no impact on 4KG5 binding. Zwick *et al.* [2003] (antibody interactions)

- A32: HIV-1 gp160deltaCT (cytoplasmic tail-deleted) proteoliposomes (PLs) containing native, trimeric envelope glycoproteins from R5 strains YU2 and JRFL, and X4 strain HXBc2, were made in a physiologic membrane setting as candidate immunogens for HIV vaccines 2F5 bound to gp160deltaCT with a reconstituted membrane ten-fold better than the same protein on beads anti-CD4BS MAbs IgG1b12 and F105, A32 (C1-C4), C11 (C1-C5), and 39F (V3) MAbs bound gp160deltaCT PLs indistinguishably from gp160deltaCT expressed on the cell surface non-neutralizing MAbs C11 and A32 bound with lower affinity than NAb IgG1b12 the MAb 17b was sCD4 inducible on gp160deltaCT PL. Grundner et al. [2002] (vaccine antigen design)
- A32: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-299) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Authors describe the epitope as N-terminal, discontinuous. Kwong et al. [2002] (antibody binding site definition and exposure)
- A32: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin stabilized oligomer gp140 delta683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and C11, A32, and 30D which did not bind the stabilized oligomer. Yang et al. [2002] (antibody binding site definition and exposure)
- A32: Using a cell-fusion system, it was found CD4i antibodies 17b, 48d, and CG10 reacted faintly with Env expressing HeLA cells even in the absence of sCD4 or CD4 expressing target cells. Reactivity increased after sCD4 addition, but not after CD4 expressing target cell addition, and binding was not increased at the cell-to-cell CD4-Env interface. This suggests the CD4i co-receptor binding domain is largely blocked at the cell-fusion interface, and so CD4i antibodies would not

- be able access this site and neutralize cell-mediated viral entry. However, CD4i MAbs 8F101 and A32, that bind outside the co-receptor domain, had a different pattern. They reacted after the formation of gp120-CD4-CXCR4 tri-complexes, so co-receptor interactions allowed exposure of their epitopes. Finnegan *et al.* [2001] (antibody binding site definition and exposure)
- A32: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang *et al.* [2000] (vaccine antigen design)
- A32: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits – a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen – SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 – nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120gp41 complexes. Binley et al. [1999] (antibody binding site definition and exposure)
- A32: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer. Binley *et al.* [1998] (antibody binding site definition and exposure)
- A32: Enhances binding of CD4i MAbs 17b and 48d, and a MAb generated in response to gp120-CD4 complex, CG10. Sullivan *et al.* [1998b] (antibody interactions)
- A32: Abs that recognize discontinuous epitopes can identify mimotopes from a phage peptide display library A32 has a unique epitope involving mostly C2 but C1 and C4 contribute six quite variable phage inserts were recognized, with a consensus of LPWYN a central Trp was the most conserved element, consistent with W427 being an important residue for binding gp120. Boots *et al.* [1997] (antibody binding site definition and exposure, mimotopes)
- A32: Review. Burton & Montefiori [1997] (review)
- A32: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric env binding A32 bound monomer, did not bind

oligomer or neutralize JRFL. Fouts *et al.* [1997] (**antibody** • C11: Of 35 Env-specific MAbs tested, only 2F5, 4E10, **binding site definition and exposure**) • C11: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G)

- A32: Does not neutralize TCLA strains or primary isolates.
   Parren et al. [1997b] (variant cross-recognition or cross-neutralization)
- A32: Binds efficiently to sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding. Wyatt et al. [1997] (antibody binding site definition and exposure)
- A32: Reciprocal inhibition of binding of anti-C1, -C5, -C4, -V3 and anti-CD4 binding site MAbs induces binding of some anti-V2 and sCD4 inducible MAbs (48d and 17b) very similar competition pattern between 2/11c, A32 and 211/c are unique among known human and rodent MAbs. Moore & Sodroski [1996] (antibody binding site definition and exposure, antibody interactions)
- A32: Does not neutralize JR-FL, or any strain strongly partial inhibition of gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola *et al.* [1996a] (co-receptor)
- A32: Not neutralizing binds domains that interact with gp41 MIP-1alpha binding to CCR-5 expressing cells can be inhibited by gp120-sCD4 and binding of A32 does not block this inhibition. Wu *et al.* [1996] (antibody binding site definition and exposure)
- A32: Review: epitope is distinct from CD4BS MAbs, 48d and 17b, and 2G12. Moore & Ho [1995] (antibody binding site definition and exposure)
- A32: Epitope is better exposed upon CD4 binding to gp120

   binding of A32 enhances binding of 48d and 17b studies using a V1/V2 deletion mutant demonstrated that enhanced binding of 48d in the presence sCD4 involves the V1/V2 loops, with more significant involvement of V2. Wyatt et al. [1995] (antibody binding site definition and exposure, antibody interactions)
- A32: Reacted with virtually every gp120 monomer of every clade tested, most conserved gp120 monomer epitope known.
   Moore et al. [1994b] (variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 1017

MAb ID C11 (c11)

**HXB2** Location Env

**Author Location** gp120

Epitope

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human

**Ab Type** gp120 C1-C5

Research Contact James Robinson, Tulane University, LA

Reference

References Selvarajah et al. 2005; Haynes et al. 2005; Pantophlet et al. 2004; Pantophlet et al. 2003b; Ohagen et al. 2003; Raja et al. 2003; Kwong et al. 2002; Basmaciogullari et al. 2002; Grundner et al. 2002; Yang et al. 2002; Binley et al. 1999; Sullivan et al. 1998b; Parren et al. 1997b; Wyatt et al. 1997; Fouts et al. 1997; Binley et al. 1997a; Wu et al. 1996; Trkola et al. 1996a; Moore & Sodroski 1996; Moore et al. 1994d; Robinson et al. 1992

**Keywords** vaccine antigen design, vaccine-specific epitope characteristics

C11: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. C11 has no indication of polyspecific autoreactivity. Haynes et al. [2005]

- C11: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. V1/V2/V3 MAb 4KG2, C1-C4 MAb A32, C1-C5 MAb C11, and HIVIG all either did not bind or had significantly diminished binding to both antigen constructs. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)
- C11: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including C11. Pantophlet *et al.* [2004]
- C11: Env genes derived from uncultured brain biopsy samples from four HIV-1 infected patients with late-stage AIDS were compared to env genes from PBMC samples. Brain isolates did not differ in the total number or positions of N-glycosylation sites, patterns of coreceptor usage, or ability to be recognized by gp160 and gp41 MAbs. C11 recognized most variants, some from each of the four individuals, by gp120 immunoprecipitation. Ohagen et al. [2003]
- C11: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional Nlinked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b]
- C11: This paper shows that binding of CD4BS MAbs to Env blocks the conformational shift that allows co-receptor CCR5 binding and CD4-independent mediated cell fusion. C11 was used as a negative control, as C11 binding did not alter binding of CD4-independent gp120 to CCR5, nor binding to CCR5expressing Cf2Th cells. Raja et al. [2003]
- C11: gp120 mutants were used to define the CXCR4 binding site using CXCR4 displayed on paramagnetic proteoliposomes (PMPLs) to reduce non-specific gp120 binding basic residues in the V3 loop and the beta19 strand (RIKQ, positions 419-422) were involved, and deletion of the V1-V2 loops allowed CD4-independent CXCR4 binding MAbs 17b (CD4i) and F105 (CD4BS) were used to study conformational changes in

the mutants – C11 was used to detect gp120 binding to CXCR4 or CCR5 on PMPLs. Basmaciogullari *et al.* [2002]

- C11: HIV-1 gp160deltaCT (cytoplasmic tail-deleted) proteoliposomes (PLs) containing native, trimeric envelope glycoproteins from R5 strains YU2 and JRFL, and X4 strain HXBc2, were made in a physiologic membrane setting as candidate immunogens for HIV vaccines 2F5 bound to gp160deltaCT with a reconstituted membrane ten-fold better than the same protein on beads anti-CD4BS MAbs IgG1b12 and F105, A32 (C1-C4), C11 (C1-C5), and 39F (V3) MAbs bound gp160deltaCT PLs indistinguishably from gp160deltaCT expressed on the cell surface non-neutralizing MAbs C11 and A32 bound with lower affinity than NAb IgG1b12 the MAb 17b was sCD4 inducible on gp160deltaCT PL. Grundner et al. [2002]
- C11: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-299) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Authors describe the epitope as N-term and C-term binding. Kwong et al. [2002]
- C11: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin stabilized oligomer gp140 delta683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and C11, A32, and 30D which did not bind the stabilized oligomer. Yang *et al.* [2002]
- C11: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 MAbs that

bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 – anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley *et al.* [1999]

- C11: Does not compete with binding of MAb generated in response to gp120-CD4 complex, CG10. Sullivan et al. [1998b]
- C11: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding C11 bound monomer, did not bind oligomer or neutralize JRFL. Fouts et al. [1997]
- C11: Does not neutralize TCLA strains or primary isolates. Parren *et al.* [1997b]
- C11: Binds efficiently to sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding partial re-exposure if sCD4 was bound does not bind to HXBc2 gp120 if the 19 C-term amino acids are deleted. Wyatt et al. [1997]
- C11: Binding enhanced by anti-V3 MAb 5G11 reciprocal inhibition with anti-C1 MAbs. Moore & Sodroski [1996]
- C11: Does not neutralize JR-FL nor block gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola *et al.* [1996a]
- C11: Did not block ability of gp120-sCD4 complexes to inhibit MIP-1alpha binding – binds to gp41-binding domain. Wu et al. [1996]
- C11: Mutations that inhibit binding: C1 (45 W/S, 88 N/P) V5 (463 N/D) and C5 (491 I/F, 493 P/K and 495 G/K) and enhance binding: C1 (36 V/L) V1-V2 (152/153 GE/SM) and DeltaV1/V2/V3. Moore *et al.* [1994d]

**No.** 1018

MAb ID L81

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 C1-C5

References Parren et al. 1997b; Ditzel et al. 1997

- L81: gp120 immobilized on solid phase by capture with anti-CD4 BS MAb L72 was used for selection of Fabs – L81 binding is abolished by C1 substitution 45 W/S, C5 substitution 491 I/F, and C3 substitution L/A. Ditzel *et al.* [1997]
- L81: Does not neutralize TCLA strains or primary isolates. Parren *et al.* [1997b]

**No.** 1019

MAb ID B2C

**HXB2 Location** Env

Author Location gp120 (HIV2ROD)

Epitope HYQ(core)

**Neutralizing** L

Immunogen vaccine

Vector/Type: peptide Strain: HIV-2 ROD

Species (Isotype) mouse

Ab Type gp120 C3

References Matsushita et al. 1995

 B2C: Viral neutralization was type-specific for HIV-2 ROD. Matsushita et al. [1995]

**No.** 1020

MAb ID polyclonal

**HXB2 Location** Env

**Author Location** 

**Epitope** 

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

Ab Type gp120 C3

References Wang et al. 2002b

 Autologous NAbs were studied in 3 patients on HAART that rebounded – phylogenetic analysis of env (V1-V5) sequences indicated that rebound viruses had evolved from or preexisted in baseline populations – HIV-1 rebound viruses from all 3 patients were resistant to neutralization by autologous IgG, unlike the baseline viruses – mutations in the C3 region was responsible for conferring neutralization resistance against autologous antibody in 2 of 3 patients. Wang et al. [2002b]

**No.** 1021

**MAb ID** 1024

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

**Immunogen** 

Species (Isotype)

Ab Type gp120 C4

References Berman et al. 1997

• 1024: Binds to 1/7 isolates from breakthrough cases from a MN gp120 vaccine trial. Berman *et al.* [1997]

**No.** 1022

MAb ID 4KG5

HXB2 Location Env

Author Location gp120 (JR-FL)

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

**Ab Type** gp120 C4, gp120 V3, gp120 V1-V2

References Selvarajah et al. 2005; Zwick et al. 2003

**Keywords** antibody binding site definition and exposure, antibody generation, antibody interactions, structure, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-

recognition or cross-neutralization

• 4KG5: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. V1/V2/V3 MAb 4KG2, C1-C4 MAb A32, C1-C5 MAb C11, and HIVIG all either did not bind or had significantly diminished binding

to both antigen constructs. Selvarajah *et al.* [2005] (vaccine antigen design, vaccine-specific epitope characteristics)

• 4KG5: 4KG5, a single-chain Fv (scFv), reacts with a conformational epitope that is formed by the V1, V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. 4KG5 was derived from the serum of HIV-1 infected patient FDA2, who showed broad neutralizing activity, but is not itself neutralizing. Denaturation of gp120 abolished binding of 4KG5 and Fab b12. Additionally, binding of 4KG5 was abrogated when any of the V1, V2 or V3 loops were deleted. Of a panel of Abs tested, only NAb b12 enhanced 4KG5 binding to gp120 JR-FL. MAbs to the following regions diminished or abrogated binding: V2 loop MAbs (G3-4, G3-136), V3 loop MAbs (19b, 447-52D, hNM01, AH48, loop2, F425 B4e8, 694-88D), V3-C4 (G3-299, G3-42, G3-519, G3-537), CD4BS (b6, b3, F91, F105, 15e, L33, 1008-D, 654-30D, 559-64D, 1027-30D, Ia3, Ia7, FG39, Fbb14). MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1, V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. 4KG5 recognized HIV-1 envelope proteins derived from JR-FL, JR-CSF, BaL, ADA and R2, but not MN, DH123, HxB2, YU2, SF2 and 89.6. Binding of 4KG5 to different strains of HIV-1 env is probably due to sequence differences in V3 and C4, rather than V1 or V2. Zwick et al. [2003] (antibody binding site definition and exposure, antibody generation, antibody interactions, variant cross-recognition or crossneutralization, structure)

**No.** 1023

MAb ID 23A (2.3A)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype)

Ab Type gp120 C5

Research Contact James Robinson, Tulane University, LA

**References** Schulke *et al.* 2002; Binley *et al.* 1999; Fouts *et al.* 1997; Trkola *et al.* 1996a; Wu *et al.* 1996; Thali *et al.* 1993; Thali *et al.* 1992a

- 23A: Ab binding characteristics of SOS gp140 were tested using SPR and RIPA SOS gp140 is gp120-gp41 bound by a disulfide bond NAbs 2G12, 2F5, IgG1b12, CD4 inducible 17b, and 19b bound to SOS gp140 better than uncleaved gp140 (gp140unc) and gp120 non-neutralizing MAbs 2.2B (binds to gp41 in gp140unc) and 23A (binds gp120) did not bind SOS gp140. Schulke *et al.* [2002]
- 23A: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and

G3-519 - nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999]

- 23A: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric env binding - 23A bound monomer, did not bind oligomer or neutralize JRFL. Fouts et al. [1997]
- 23A: C5 binding MAb does not inhibit gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]
- 23A: Called 2.3A Did not block ability of gp120-sCD4 complexes to inhibit MIP-1alpha binding - binds to gp41-binding domain of gp120. Wu et al. [1996]

No. 1024

**MAb ID** D7324

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen vaccine

HIV component: gp120

Species (Isotype) sheep

Ab Type gp120 C5

Research Contact Aalto BioReagents Ltd, Dublin, Ireland or Cliniqa Inc., Fallbrook, CA, USA

References Martín-García et al. 2005; Koefoed et al. 2005; Jeffs et al. 2004; Zwick et al. 2003; Herrera et al. 2003; Poignard et al. 2003; Basmaciogullari et al. 2002; Xiang et al. 2002a; Gram et al. 2002; Sanders et al. 2002; Binley et al. 1998; Mondor et al. 1998; Ugolini et al. 1997; Ditzel et al. 1997; Trkola et al. 1996a; Wyatt et al. 1995; Moore et al. 1993b; Moore et al. 1993a; Sattentau & Moore 1991; Moore

Keywords antibody binding site definition and exposure, antibody interactions, subtype comparisons, vaccine antigen design

- D7324: IgG antibody phage display libraries were created from HIV-1 + individuals after pre-selection of PBMC with gp120, as an alternative to using bone marrow for generating libraries. D7324 was among a set of Abs used for competition studies to define the binding sites of the newly isolated MAbs, representing a MAb with a C5 epitope. Koefoed et al. [2005] (antibody binding site definition and exposure)
- D7324: The HIV-1 Bori-15 variant was adapted from the Bori isolate for replication microglial cells. Bori-15 had increased replication in microglial cells and a robust syncytium-forming phenotype, ability to use low levels of CD4 for infection, and increased sensitivity to neutralization by sCD4 and 17b. Four amino acid changes in gp120 V1-V2 were responsible for this change. Protein functionality and integrity of soluble, monomeric gp120-molecules derived from parental HIV-1 Bori and microglia-adapted HIV-1 Bori-15 was assessed in ELISA

binding assays using F105, IgG1b12, 17b and 48d, 2G12 and 447-52D. Association rates of sCD4 and 17b were not changed. but dissociation rates were 3-fold slower for sCD4 and 14-fold slower for 17b. The kinetics of D7324 binding were tested as a control, and were unchanged. Martín-García et al. [2005] (antibody binding site definition and exposure)

- D7324: A set of oligomeric envelope proteins were made from six primary isolates for potenial use as vaccine antigens: 92/UG/037 (clade A), HAN2/2 (clade B), 92/BR25/025 (clade C), 92/UG/021 (clade D), 93/BR/029 (clade F) and MVP5180 (clade O). This was one of a panel of MAbs used to explore folding and exposure of well characterized epitopes. The clade C isolate BR25 is apparently misfolded, as conformationdependent antibodies did not bind to it. D7324 bound to clade A, B, C, D and F HIV-1 primary isolates, but not to the group O protein. Polyclonal sera raised in rabbits against these antigens cross-bound the other antigens, but none of the sera had neutralizing activity. Jeffs et al. [2004] (vaccine antigen design, subtype comparisons)
- D7324: Used to capture gp120 onto solid phase for epitope mapping. Basmaciogullari et al. [2002]; Binley et al. [1998]; Ditzel et al. [1997]; Herrera et al. [2003]; Moore et al. [1993a,b]; Poignard et al. [2003]; Sanders et al. [2002]; Xiang et al. [2002a]
- D7324: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a C5-binding polyclonal Ab that had no impact on 4KG5 binding. Zwick et al. [2003] (antibody interactions)
- D7324: Called NEA9205 gp120 capture ELISAs with MAbs D7324 (anti-C-term) or 9205 (anti-V3) were compared in a study of orientation of glycosylation sites - CD4 binding could only inhibit deglycosylation when gp120 was bound to the plate by D7324, not by 9205, while Abs from HIV-1 infected people inhibited deglycosylation most effectively when gp120 was caught by 9205. Gram et al. [2002]
- D7324: Epitope in C5 Does not neutralize JR-FL nor block gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a]
- D7324: Binds to the last 15 amino acids in gp120 used for antigen capture ELISA. Wyatt et al. [1995]
- D7324: Binding unaltered by gp120 binding to sCD4, in contrast to 110.5, 9284, 50-69 and 98-6. Sattentau & Moore [1991]

**No.** 1025

MAb ID 10/46c

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component:

gp120

Species (Isotype) rat

**Ab Type** gp120 CD4BS

References Peet et al. 1998; Jeffs et al. 1996; Cordell et al. • 1027-30-D: Called 1027-30D. scFv 4KG5 reacts with a con-

- 10/46c: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic – these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind – 10/46c was not affected by V3 serine substitutions – mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]
- 10/46c: Increased binding when V1/V2 or V1/V2 and V3 were deleted from gp120. Jeffs et al. [1996]

**No.** 1026

**MAb ID** 1008-D

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

Research Contact Susan Zolla-Pazner (Zol-Med

las01@mcrcr6.med.nyu), NYU

Center, NY, NY

References Zwick et al. 2003; Zolla-Pazner et al. 1995

Keywords antibody interactions

• scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. Research Contact Shermaine Tilley, Public Health Research In-MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick et al. [2003] (antibody interactions)

**No.** 1027

**MAb ID** 1027-30-D (1027-30D)

**HXB2 Location** Env

Author Location Env

**Epitope** 

**Neutralizing** 

**Immunogen** 

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 CD4BS

Research Contact Susan Zolla-Pazner (Zol-

> las01@mcrcr6.med.nyu) (NYU Med.

Center)

References Gorny & Zolla-Pazner 2004; Zwick et al. 2003; Hioe et al. 2000

**Keywords** antibody interactions, review

• 1027-30D: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)

- formational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick et al. [2003] (antibody interactions)
- 1027-30-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses - anti-CD4 binding site MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells - CD4BS MAbs 654-D, 559/64-D, 588-D, 830-D, 1027-30-D, and 1202-30D strongly diminished proliferation. Hioe et al. [2000]

No. 1028

MAb ID 1125H (1125h)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L (MN)

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type gp120 CD4BS

stitute, USA

References Yang et al. 1998; Alsmadi & Tilley 1998; Wyatt et al. 1998; Pincus et al. 1996; Warrier et al. 1996; D'Souza et al. 1995; Pinter et al. 1993b; Wyatt et al. 1992; Thali et al. 1992a; Tilley et al. 1991a; Tilley et al. 1991b

- 1125H: A study of 6 anti-Env MAbs and their ability to bind or direct ADCC against target cells infected with IIIB, MN, SF-2, and RF – bound and directed lysis against all four strains. Alsmadi & Tilley [1998]
- 1125H: Called 1125h summary of the implications of the crystal structure of the core of gp120 bound to CD4 and 17b with what is known about mutations that reduce NAb binding probable mechanism of neutralization by CD4BS Ab is direct interference with CD4 binding. Wyatt et al. [1998]
- 1125H: A neutralization assay was developed based on heminested PCR amplification of the LTR (HNPCR) - LTR-HNPCR consistently revealed HIV DNA and was shown to be a rapid, specific and reliable neutralization assay based on tests with 6 MAbs and 5 isolates. Yang et al. [1998]
- 1125H: A panel of immunotoxins were generated by linking Env MAbs to ricin A – immunotoxins mediated cell killing, but killing was not directly proportional to binding. Pincus et al. [1996]
- 1125H: Synergistic neutralization of HIV-1 when combined with anti-V2 MAb C108G. Warrier et al. [1996]
- 1125H: Neutralization was MN specific failed to neutralize JRCSF, and 2 B subtype and 1 D subtype primary isolates in a multi-laboratory study involving 11 labs. D'Souza et al. [1995]

- an anti-V3 HuMAb, 41148D. Pinter et al. [1993b]
- 1125H: Amino acid substitutions in HXB2 that strongly inhibit binding: 88, 102, 117, 113, 257, 368, 370, 421, 427, 457, 470, 480. Thali et al. [1992a]
- 1125H: Precipitation of Delta 297-329 env glycoprotein, with has a deleted V3 loop, is much more efficient that precipitation of wild type. Wyatt et al. [1992]
- 1125H: Binding to gp120 inhibited by CD4 epitope is destroyed by reduction, but not by removal of N-linked sugars potent neutralization of MN, RF, SF-2 and IIIB - neutralization synergy with anti-V3 MAb 4117C. Tilley et al. [1991a]

No. 1029

**MAb ID** 1125H (1125h)

**HXB2 Location** Env Author Location gp120

**Epitope** 

Subtype B

**Neutralizing** L (MN)

Immunogen HIV-1 infection

Species (Isotype) human (IgG1 $\kappa$ )

**Ab Type** gp120 CD4BS

Research Contact Shermaine Tilley, Public Health Research Institute, USA

> References Pinter et al. 2004; Gorny & Zolla-Pazner 2004; Yang et al. 1998; Alsmadi & Tilley 1998; Wyatt et al. 1998; Pincus et al. 1996; Warrier et al. 1996; D'Souza et al. 1995; Pinter et al. 1993b; Wyatt et al. 1992; Thali et al. 1992a; Tilley et al. 1991a; Tilley et al. 1991b

Keywords ADCC, antibody binding site definition and exposure, antibody interactions, assay development, immunotoxin, review, structure, subtype comparisons, variant cross-recognition or cross-neutralization

- 1125H: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 1125H: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Three anti-CD4BS MAbs were tested, including IgG1b12 which neutralizes both JRFL and SF162. The affinities for IgG1b12 and 5145A were similar for both JRFL and SF612, but 1125A bound with 2.5 fold higher affinity to SF162. 5145A and 1125H both preferentially neutralize SF162, but not JRFL, and Research Contact Virus Testing Systems Corp., Houston, TX the CD4BS is more sensitive to neutralization in the context of the SF162 V1V2 loop. This was also true for neutralization by sCD4. Pinter et al. [2004] (variant cross-recognition or cross-neutralization)

• 1125H: Binding to soluble gp120 enhanced by the presence of • 1125H: A study of 6 anti-Env MAbs and their ability to bind or direct ADCC against target cells infected with IIIB, MN, SF-2, and RF – bound and directed lysis against all four strains. Alsmadi & Tilley [1998] (ADCC)

- 1125H: Called 1125h summary of the implications of the crystal structure of the core of gp120 bound to CD4 and 17b with what is known about mutations that reduce NAb binding probable mechanism of neutralization by CD4BS Ab is direct interference with CD4 binding. Wyatt et al. [1998] (structure)
- 1125H: A neutralization assay was developed based on heminested PCR amplification of the LTR (HNPCR) - LTR-HNPCR consistently revealed HIV DNA and was shown to be a rapid, specific and reliable neutralization assay based on tests with 6 MAbs and 5 isolates. Yang et al. [1998] (assay development)
- 1125H: A panel of immunotoxins were generated by linking Env MAbs to ricin A – immunotoxins mediated cell killing, but killing was not directly proportional to binding. Pincus et al. [1996] (**immunotoxin**)
- 1125H: Synergistic neutralization of HIV-1 when combined with anti-V2 MAb C108G. Warrier et al. [1996] (antibody interactions)
- 1125H: Neutralization was MN specific failed to neutralize JRCSF, and 2 B subtype and 1 D subtype primary isolates in a multi-laboratory study involving 11 labs. D'Souza et al. [1995] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 1125H: Binding to soluble gp120 enhanced by the presence of an anti-V3 HuMAb, 41148D. Pinter et al. [1993b] (antibody interactions)
- 1125H: Amino acid substitutions in HXB2 that strongly inhibit binding: 88, 102, 117, 113, 257, 368, 370, 421, 427, 457, 470, 480. Thali et al. [1992a] (antibody binding site definition and exposure)
- 1125H: Precipitation of Delta 297-329 env glycoprotein, with has a deleted V3 loop, is much more efficient that precipitation of wild type. Wyatt et al. [1992] (antibody binding site definition and exposure)
- 1125H: Binding to gp120 inhibited by CD4 epitope is destroyed by reduction, but not by removal of N-linked sugars neutralization of MN, RF, SF-2 and IIIB - neutralization synergy with anti-V3 MAb 4117C. Tilley et al. [1991a] (antibody binding site definition and exposure, antibody interactions, variant cross-recognition or cross-neutralization)

**No.** 1030

MAb ID 120-1B1

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

**Immunogen** 

Species (Isotype) human

Ab Type gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Watkins et al.

**Keywords** antibody binding site definition and exposure,

(Zol-

• 120-1B1: This review summarizes MAbs directed to HIV-1 Research Contact Susan Env. There are 51 CD4BS MAbs and Fabs in the database; las016 most, like this MAb, neutralize TCLA strains only. Gorny & Center Zolla-Pazner [2004] (review) References Gorny

120-1B1: A neutralization escape mutant (HXB2 A281V) was selected by growth of HXB2 in the presence of broadly neutralizing sera – 120-1B1 was not affected by this mutation. Watkins *et al.* [1993] (antibody binding site definition and exposure)

No. 1031

MAb ID 1202-D (1202-30-D)

**HXB2 Location** Env

**Author Location** Env

Epitope

Neutralizing

**Immunogen** 

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 CD4BS

Research Contact Susan Zolla-Pazner

las01@mcrcr6.med.nyu) (NYU Med.

Center)

**References** Gorny & Zolla-Pazner 2004; Nyambi *et al.* 2000; Hioe *et al.* 2000; Nyambi *et al.* 1998

Keywords review, subtype comparisons

- 1202-D: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 1202-D: Called 1202-30D Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses anti-CD4 binding site MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells CD4BS MAbs 654-D, 559/64-D, 588-D, 830-D, 1027-30-D, and 1202-30D strongly diminished proliferation. Hioe *et al.* [2000]
- 1202-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 CD4BS MAbs – CD4BS MAbs bound consistently to most isolates of clade D, but poorly to isolates of other clades with the exception of broadly reactive MAb IgG1b12. Nyambi et al. [2000] (subtype comparisons)
- 1202-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H CD4-BS Abs tended to bind weakly without clade specificity to virions, but bound well to soluble gp120 1202-D did not bind to any B clade viruses, and weakly bound A, C, and G clade isolates 559/64-D, 558-D and 1202-D had similar reactivities. Nyambi *et al.* [1998] (subtype comparisons)

**No.** 1032

**MAb ID** 1331E

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope

Neutralizing

**Immunogen** HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 CD4BS

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med. Center)

**References** Gorny & Zolla-Pazner 2004; Gorny *et al.* 2000

**Keywords** antibody binding site definition and exposure, review

- 1331E: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 1331E: Inhibits sCD4 binding to rec gp120 LAI binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer CD4BS MAbs 559/64-D, 654-D, 729-D, 9CL and 1331E bound with a 5-13 fold preference for the oligomer. Gorny *et al.* [2000] (antibody binding site definition and exposure)

**No.** 1033

**MAb ID** 1570 (1570A, 1570C, 1570D)

**HXB2** Location Env

Author Location Env (PR12, BH10)

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Jeffs *et al.* 2001 Keywords antibody binding site definition and exposure, antibody generation, review, subtype comparisons, variant cross-recognition or cross-neutralization

- 1570: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 1570: BH10 was mutated to form the PR12 protein with the first 74 C-terminal amino acids and the V1, V2 and V3 hypervariable loops deleted and a very well exposed CD4 binding domain (CD4bd) this proteins was used to select three new human CD4BS MAbs 1570, 1595 and 1599 three MAbs were isolated from one individual, 1570A, C and D but all were determined to have the same V(H)3 region 1570 was able to bind to a panel of recombinant proteins from the A, B, C, D, and E subtypes. Jeffs et al. [2001] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 1034

**MAb ID** 1595

**HXB2 Location** Env

Author Location Env (PR12, BH10)

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Jeffs et al. 2001 Keywords antibody binding site definition and exposure, antibody generation, review, subtype comparisons, variant cross-recognition or crossneutralization

- 1595: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 1595: BH10 was mutated to form the PR12 protein with the first 74 C-terminal amino acids and the V1, V2 and V3 hypervariable loops deleted and a very well exposed CD4 binding domain (CD4bd) - this proteins was used to select three new human CD4BS MAbs 1570, 1595 and 1599 – 1595 was able to bind gp120 from the A, B, and D clades from a panel of recombinant proteins from the A, B, C, D, and E subtypes. Jeffs et al. [2001] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

No. 1035

**MAb ID** 1599

**HXB2 Location** Env

Author Location Env (PR12, BH10)

**Epitope Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Jeffs et al. 2001 Keywords antibody binding site definition and exposure, • 15e: UK Medical Research Council AIDS reagent: ARP3016.

antibody generation, review, subtype comparisons, variant cross-recognition or crossneutralization

- 1599: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 1599: BH10 was mutated to form the PR12 protein with the first 74 C-terminal amino acids and the V1, V2 and V3 hypervariable loops deleted and a very well exposed CD4 binding domain (CD4bd) - this proteins was used to select three new human CD4BS MAbs 1570, 1595 and 1599 - 1599 was able to bind gp120 only from the B clade from a panel of recombinant proteins from the A, B, C, D, and E subtypes. Jeffs et al. [2001] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 1036

**MAb ID** 15e (1.5e, 1.5E, 15E)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

**Ab Type** gp120 CD4BS

Research Contact James Robinson, Tulane University, LA, and

David Ho, ADARC, NY, NY

References Nabatov et al. 2004; Pantophlet et al. 2004;

Gorny & Zolla-Pazner 2004: Pantophlet et al. 2003b; Zwick et al. 2003; Raja et al. 2003; Pantophlet et al. 2003a; Kwong et al. 2002; Zhang et al. 2002; Xiang et al. 2002b; Kolchinsky et al. 2001; Park et al. 2000; Sullivan et al. 1998a; Fouts et al. 1998; Trkola et al. 1998; Binley et al. 1998; Sullivan et al. 1998b; Parren et al. 1998a; Wyatt et al. 1998; Parren et al. 1997b; Berman et al. 1997; Wyatt et al. 1997; Li et al. 1997; Fouts et al. 1997; Binley et al. 1997a; Wisnewski et al. 1996; McDougal et al. 1996; Trkola et al. 1996a; Poignard et al. 1996a; Moore & Sodroski 1996; McKeating et al. 1996; Lee et al. 1995; Sattentau & Moore 1995; Moore et al. 1994a; Moore et al. 1994b; Cook et al. 1994; Thali et al. 1994; Bagley et al. 1994; Wyatt et al. 1993; Watkins et al. 1993; Thali et al. 1993; Moore & Ho 1993; Takeda et al. 1992; Thali et al. 1992a; Wyatt et al. 1992; Ho et al. 1992; Koup et al. 1991; Ho et al. 1991b; Cordell et al. 1991; Thali et al. 1991; Robinson et al. 1990a

Keywords ADCC, adjuvant comparison, antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain, brain/CSF, co-receptor, enhancing activity, review, structure, subtype comparisons, vaccine antigen design, variant crossrecognition or cross-neutralization

- 15e: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 15e: A set of HIV-1 chimeras that altered V3 net charge and glycosylation patterns in V1V2 and V3, involving inserting V1V2 loops from a late stage primary isolate taken after the R5 to X4 switch, were studied with regard to phenotype, coreceptor usage, and MAb neutralization. The loops were cloned into a HXB2 envelope with a LAI viral backbone. It was observed that the addition of the late-stage isolate V1V2 region and the loss of V3-linked glycosylation site in the context of high positive charge gave an X4 phenotype. R5X4, R5, and X4 viruses were generated, and sCD4, 2G12 and b12 neutralization resistance patterns were modified by addition of the late stage V1V2, glycosylation changes, and charge in concert, while neutralization by 2F5 was unaffected. 15e, 17b, and 48d could not neutralize any of the variants tested. Nabatov et al. [2004] (antibody binding site definition and exposure, co-receptor)
- 15e: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and

- does not bind to 21 other MAbs to 7 epitopes on gp120, including 15e. Pantophlet *et al.* [2004] (vaccine antigen design)
- 15e: Alanine scanning mutagenesis was used to compare substitutions that affected anti-CD4BS NAb b12 rec gp120s were engineered to contain combinations of Alanine substitutions that enhanced b12 binding, and while binding of b12 to these gp120 monomers was generally maintained or increased, binding by five non-neutralizing anti-CD4bs MAbs (b3, b6, F105, 15e, and F91) was reduced or completely abolished. Pantophlet *et al.* [2003a] (antibody binding site definition and exposure)
- 15e: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of non-neutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet *et al.* [2003b] (vaccine antigen design)
- 15e: This paper shows that binding of CD4BS MAbs to Env blocks the conformational shift that allows co-receptor CCR5 binding and CD4-independent mediated cell fusion. CD4BS MAbs F105, 15e, and IgG1b12 as well as their Fab fragments inhibited CD4-independent binding of the V1/V2 loop-deleted gp120 glycoproteins of R5 HIV-1 isolates ADA, YU2 and JRFL and to CCR5 in a concentration dependent manner. Raja et al. [2003] (co-receptor)
- 15e: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick et al. [2003] (antibody interactions)
- 15e: Called 1.5e. Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar. Enthalpy and entropy changes were divergent, but compensated. Not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs (17b, 48d, 1.5e, b6, F105 and F91) had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, but the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol, The high values suggest surface burial or protein folding an ordering of amino acids. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab

binding not faced by other anti-gp120 Abs. Kwong *et al.* [2002] (antibody binding site definition and exposure)

- 15e: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations 375 S/W seems to favor a conformation of gp120 closer to the CD4-bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang et al. [2002b] (antibody binding site definition and exposure)
- 15e: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002] (antibody binding site definition and exposure)
- 15e: Mutations in two glcosylation sites in the V2 region of HIV-1 ADA at positions 190 and 197 (187 DNTSYRLINCNTS 199) cause the virus to become CD4-independent and able to enter cells through CCR5 alone these same mutations tended to increase the neutralization sensitivity of the virus, including to 15e. Kolchinsky *et al.* [2001] (antibody binding site definition and exposure)
- 15e: Six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park *et al.* [2000] (antibody binding site definition and exposure)
- 15e: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer CD4BS MAbs 15e, F91 and IgG1b12 bound better to the deleted protein than to wild type. Binley *et al.* [1998] (antibody binding site definition and exposure)
- 15e: CD4BS MAbs 15e, 21h, and F91 bind with even lower affinity than 205-43-1 and 205-42-15 to JRFL oligomer. Fouts et al. [1998] (antibody binding site definition and exposure)
- 15e: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren *et al.* [1998a] (antibody binding site definition and exposure)

- 15e: Competes with CG-10 binding, a MAb raised against a gp120 CD4 complex, this was probably due to the disruption of CD4-gp120 by 15e. Sullivan *et al.* [1998b] (antibody binding site definition and exposure, antibody interactions)
- 15e: Called 1.5e the HIV-1 virus YU2 entry can be enhanced by MAbs binding to the CD4BS, V3 loop, and CD4i epitopes the activation for this enhanced entry state could be conferred on HxB2 by introducing the YU2 V3 loop, or the YU2 V3 and V1/V2 loops a similar effect is observed by sub-neutralizing concentrations of sCD4 and the effect is dependent of CCR5 1.5e enhances and does not neutralize YU2 env even at 50 ug/ml. Sullivan *et al.* [1998a] (antibody binding site definition and exposure)
- 15e: No detectable neutralizing activity among primary isolates with different co-receptor usage – some neutralization of TCLA strains. Trkola *et al.* [1998] (variant cross-recognition or cross-neutralization)
- 15e: Summary of the implications of the crystal structure of the core of gp120 bound to CD4 and 17b with what is known about mutations that reduce NAb binding probable mechanism of neutralization by CD4BS Ab is direct interference with CD4 binding. Wyatt et al. [1998] (structure)
- 15e: Called 1.5E Binds to 7/7 isolates from breakthrough cases from a MN gp120 vaccine trial. Berman *et al.* [1997] (variant cross-recognition or cross-neutralization)
- 15e: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding 15e bound monomer, did not bind oligomer or neutralize JRFL. Fouts *et al.* [1997] (antibody binding site definition and exposure)
- 15e: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env 15e could only achieve 50% neutralization, but could act synergistically with anti-V3 MAb 694/98-D to achieve 90%. Li *et al.* [1997] (antibody interactions)
- 15e: Neutralizes TCLA strains, but not primary isolates. Parren *et al.* [1997b]
- 15e: Does not bind to HXBc2 gp120 if the 19 C-term amino acids, in conjunction with C1 positions 31-93, are deleted. Wyatt et al. [1997] (antibody binding site definition and exposure)
- 15e: Neutralizes HIV-1 LAI less potently than V3 specific MAbs. McDougal *et al.* [1996]
- 15e: Called 1.5e Neutralizes HXB2, but fails to neutralize chimeric virus with gp120 from primary isolates in an HXB2 background. McKeating *et al.* [1996] (variant cross-recognition or cross-neutralization)
- 15e: gp120 binding enhanced by anti-V3 MAb 5G11 and anti-V2 MAb G3-136 binding inhibited by other CD4 binding site MAbs, antibodies that bind to gp120 only when CD4 is bound, and CD4-IgG. Moore & Sodroski [1996] (antibody interactions)
- 15e: Anti-CD4BS MAbs 15e, 21h, and IgG1b12 did not cause gp120 dissociation from virus, or exposure of the gp41 epitope of MAb 50-69, in contrast to CD4i MAb 48d and anti-V3 neutralizing MAbs. Poignard *et al.* [1996a] (antibody interactions)

- 15e: Inhibits gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola *et al.* [1996a] (antibody binding site definition and exposure)
- 15e: 15e is V H4 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski *et al.* [1996] (antibody sequence, variable domain)
- 15e: The V4 and V5 domains are essential for 1.5e binding, in contrast to the V1, V2, and V3 loops. Lee *et al.* [1995] (antibody binding site definition and exposure)
- 15e: Binds with higher affinity to monomer than to oligomer, moderate association rate. Sattentau & Moore [1995] (antibody binding site definition and exposure)
- 15e: Heavy chain is V HIV, V2-1 light chain is V\_kappaI, Hum01/012. Compared to 21h and F105. Bagley et al. [1994] (antibody sequence, variable domain)
- 15e: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon anti-CD4 MAbs moderately inhibit gp120 binding to GalCer, possibly through steric hindrance binding of GalCer to gp120 inhibited but did not completely block 15e binding. Cook et al. [1994] (antibody binding site definition and exposure, brain/CSF)
- 15e: Cross-reactive with gp120 proteins from clades B and D, less so with A and C, and not reactive with clade E and F. Moore *et al.* [1994b] (subtype comparisons)
- 15e: A mutation in gp41, 582 A/T, confers resistance to neutralization (also confers resistance to MAbs F105, 48d, 21h and 17b). Thali *et al.* [1994] (antibody binding site definition and exposure)
- 15e: Conformational, does not bind denatured gp120 neutralizes IIIB reactive with SF-2 gp120 strong inhibition of HIV+ human sera binding to IIIB gp120. Moore & Ho [1993] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 15e: Called 15E a neutralization escape mutant (HXB2 A281V) was selected by growth of HXB2 in the presence of broadly neutralizing sera 15E neutralization was not affected by this mutation. Watkins *et al.* [1993] (antibody binding site definition and exposure)
- 15e: Binding to Delta V1/2 and Delta V1/2/3 mutant glycoproteins is greater than binding to wildtype gp120. Wyatt *et al.* [1993] (antibody binding site definition and exposure)
- 15e: gp120 mutants that affect 15e epitope binding: 113, 257, 368, 370, 421, 427, 475 four of these coincide with amino acids important for the CD4 binding domain. Ho *et al.* [1992] (antibody binding site definition and exposure)
- 15e: Amino acid substitutions in HXB2 that strongly inhibit binding, similar to Ho *et al.* [1992], some additional, 88, 102, 117, 113, 257, 368, 370, 421, 427, 457, 470, 480 Thali *et al.* [1992a]. Ho *et al.* [1992]; Thali *et al.* [1992a] (antibody binding site definition and exposure)
- 15e: Called N70-1.5e does not enhance infection of HIV-1 IIIB and MN. Thali *et al.* [1992a] (**enhancing activity**)
- 15e: Precipitation of Delta 297-329 env glycoprotein, with a deleted V3 loop, is much more efficient that precipitation of wild type. Wyatt et al. [1992] (antibody binding site definition and exposure)

- 15e: Cross-competes with MAbs ICR 39.13g and ICR 39.3b. 21h: CD4BS MAbs 15e, 21h, and F91 bind with even lower Cordell et al. [1991] (antibody interactions)
- 15e: Broadly neutralizing, binds multiple strains, competes with CD4 for gp120 binding, DTT reduction of env abrogates binding - more potent blocking of gp120-sCD4 binding than MAbs G3-536 and G3-537.. Ho et al. [1991b] (adjuvant comparison, variant cross-recognition or cross-neutralization)
- 15e: Binds to gp120 of HIV-1 IIIB, but not RF mediates ADCC – deletion of the V3 loop from gp120 does not alter ADCC activity. Koup et al. [1991] (ADCC, variant crossrecognition or cross-neutralization)

**No.** 1037

**MAb ID** 21h (2.1H)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 CD4BS

Research Contact James Robinson, Tulane University, LA

References Gorny & Zolla-Pazner 2004; Xiang et al. 2002b; Fouts et al. 1998; Parren et al. 1998a; Wyatt et al. 1998; Parren et al. 1997b; Wyatt et al. 1997; Ugolini et al. 1997; Li et al. 1997; Fouts et al. 1997; Binley et al. 1997a; McKeating et al. 1996; Wisnewski et al. 1996; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; Thali et al. 1994; Bagley et al. 1994; Moore et al. 1994a; Moore et al. 1994b; Moore & Ho 1993; Wyatt et al. 1993; Ho et al. 1992; Thali et al. 1992a; Ho et al. 1991b

Keywords acute/early infection, antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain, binding affinity, review, structure, subtype comparisons, variant cross-recognition or crossneutralization

- 21h: UK Medical Research Council AIDS reagent: ARP3017.
- 21h: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 21h: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations—375 S/W seems to favor a conformation of gp120 closer to the CD4bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91) was markedly reduced— IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced—2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang et al. [2002b] (antibody binding site definition and exposure)

affinity than 205-43-1 and 205-42-15 to JRFL oligomer - conclusions of this paper contrast with Parren et al. [1998a] Fouts et al. [1998]. Fouts et al. [1998]; Parren et al. [1998a] (binding affinity)

- 21h: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated – authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a] (antibody binding site definition and exposure)
- 21h: Summary of the implications of the crystal structure of the core of gp120 bound to CD4 and 17b with what is known about mutations that reduce NAb binding - probable mechanism of neutralization by CD4BS Ab is direct interference with CD4 binding. Wyatt et al. [1998] (antibody binding site definition and exposure, structure)
- 21h: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding - 21h bound monomer, did not bind oligomer or neutralize JRFL. Fouts et al. [1997] (antibody binding site definition and exposure)
- 21h: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env - 50% neutralization could not be achieved at a maximal concentration of 67 mug/ml. Li et al. [1997] (variant cross-recognition or cross-neutralization)
- 21h: Neutralizes TCLA strains, but not primary isolates. Parren et al. [1997b] (variant cross-recognition or crossneutralization)
- 21h: Viral binding inhibition by 21h strongly correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5). Ugolini et al. [1997] (antibody binding site definition and exposure)
- 21h: Binds both gp120 and soluble gp120+gp41 complex efficiently, suggesting its gp120 epitope is not blocked by gp41 binding - major deletions in C1 and C5 and deletions of the V1V2 and V3 loops do not diminish binding. Wyatt et al. [1997] (antibody binding site definition and exposure)
- 21h: Called 2.1H Neutralizes HXB2, but fails to neutralize chimeric virus with gp120 from primary isolates in an HXB2 background. McKeating et al. [1996] (variant crossrecognition or cross-neutralization)
- 21h: Anti-CD4 binding site MAb reciprocal inhibition by anti-C1, -C4 and other anti-CD4 binding site antibodies - enhanced by some anti-V2 MAbs and anti-V3 MAb 5G11 - enhances binding of some anti-V3 and -V2 MAbs. Moore & Sodroski [1996] (antibody interactions)
- 21h: Anti-CD4BS MAbs 15e, 21h, and IgG1b12 did not cause gp120 dissociation from virus, or exposure of the gp41 epitope of MAb 50-69, in contrast to CD4i MAb 48d and anti-V3 neutralizing MAbs. Poignard et al. [1996a] (antibody binding site definition and exposure, antibody interactions)
- 21h: 21h is V H3 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996] (antibody sequence, variable domain)
- 21h: Binds with higher affinity to monomer than to oligomer, moderate association rate. Sattentau & Moore [1995] (antibody binding site definition and exposure)

- 21h: Heavy chain is V HIII, VDP-35 light chain is V\_lambdaIIIa, Hum318. Compared to 15e and F105. Bagley et al. [1994] (antibody sequence, variable domain)
- 21h: Has strong cross-reactivity with gp120 monomers from most subtypes, A-F, with the least reactivity to clade E. Moore *et al.* [1994b] (**subtype comparisons**)
- 21h: Competition studies with human sera from seroconverting individuals showed that anti-CD4 BS antibodies can arise very early in infection, comparable or prior to anti-V3 antibodies. Moore *et al.* [1994a] (acute/early infection)
- 21h: A mutation in gp41, 582 A/T, confers resistance to neutralization (also confers resistance to MAbs F105, 48d, 15e and 17b). Thali *et al.* [1994] (variant cross-recognition or cross-neutralization)
- 21h: Conformational, does not bind denatured gp120 neutralizes IIIB reactive with SF-2 gp120 strong inhibition of HIV+ human sera binding to IIIB gp120. Moore & Ho [1993] (antibody binding site definition and exposure)
- 21h: Binding to Delta V1/2 and Delta V1/2/3 mutant glycoproteins is greater than binding to wildtype gp120. Wyatt *et al.* [1993] (antibody binding site definition and exposure)
- 21h: Amino acid substitutions in HXB2 that inhibit binding, some shared with CD4 binding inhibition, 88, 113, 257, 368, 370, 421, 470, 480. Thali *et al.* [1992a] (antibody binding site definition and exposure)

**No.** 1038

MAb ID 28A11/B1

**HXB2 Location** Env

Author Location gp120 (SF162)

Epitope
Subtype B
Neutralizing L
Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

 $juvant\ (MPL+TDM)\ (RIBI)$ 

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 CD4BS

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

**References** Gorny & Zolla-Pazner 2004; He *et al.* 2002 **Keywords** review, subtype comparisons, variant cross-recognition or cross-neutralization

- 28A11/B1: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 28A11/B1: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2 $\kappa$  MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—6 anti-CD4BS MAbs competed with anti-CD4BS MAb 5145A, blocked sCD4 binding and were conformationally sensitive—4/6 could neutralize the autologous strain SF162, and were broadly cross-reactive with B clade R5 and X4 strains (not E clade)—28A11/B1 was one of these four MAbs. He *et al.* [2002] (variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 1039

MAb ID 2G6

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen

Species (Isotype)

Ab Type gp120 CD4BS

Research Contact Herman Katinger, Inst. Appl. Microbiol. Uni-

versity of Agricultural Science, or Polymun

Scientific Inc., Vienna, Austria

References Gorny & Zolla-Pazner 2004; Parren et al.

1998a; Fouts et al. 1998

**Keywords** antibody binding site definition and exposure, review, variant cross-recognition or cross-

neutralization

- 2G6: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 2G6: Binds to JRFL oligomer with an affinity comparable to IgG1b12, but does not neutralize the virus, so binding of oligomer is not always predictive of neutralization conclusions of this paper contrast with Parren *et al.* [1998a] authors propose a model where 205-46-9 and 2G6 may inhibit CD4 binding, but cause a conformational shift which enhances CCR5 binding and thus counteracts the neutralizing effect Fouts *et al.* [1998]. Fouts *et al.* [1998]; Parren *et al.* [1998a] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)

**No.** 1040

MAb ID 35F3/E2

**HXB2 Location** Env

Author Location gp120 (SF162)

Epitope Subtype B

Neutralizing L

Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 CD4BS

Env. There are 51 CD4BS MAbs and Fabs in the database; **Research Contact** Dr. Abraham Pinter, Public Health Research most, like this MAb, neutralize TCLA strains only. Gorny & Institute, Newark, NJ, pinter@phri.org

**References** Gorny & Zolla-Pazner 2004; He *et al.* 2002 **Keywords** review, subtype comparisons, variant cross-

recognition or cross-neutralization

- 35F3/E2: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 35F3/E2: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2κ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—6 anti-CD4BS MAbs competed with anti-CD4BS MAb 5145A, blocked sCD4 binding and were conformationally

sensitive—4/6 could neutralize the autologous strain SF162, Species (Isotype) human (IgG1 $\lambda$ ) and were broadly cross-reactive with B clade R5 and X4 strains (not E clade)—35F3/E2 was one of these four MAbs. He et al. Research Contact Susan [2002] (variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 1041 MAb ID 38G3/A9 **HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** Subtype B **Neutralizing** L Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 CD4BS

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002 Keywords variant cross-recognition or crossneutralization

- 38G3/A9: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (variant cross-recognition or crossneutralization)
- 38G3/A9: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human  $IgG2\kappa$ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120-6 anti-CD4BS MAbs competed with anti-CD4BS MAb 5145A, blocked sCD4 binding and were conformationally sensitive—4/6 could neutralize the autologous strain SF162, and were broadly cross-reactive with B clade R5 and X4 strains (not E clade)—38G3/A9 was one of these four MAbs. He et al. [2002] (variant cross-recognition or cross-neutralization)

**No.** 1042

**MAb ID** 428

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

References Jeffs et al. 1996; Karwowska et al. 1992a

• 428: Slight, not significant increased binding when V1/V2 or V1/V2 and V3 were deleted from gp120. Jeffs et al. [1996]

**No.** 1043

**MAb ID** 448-D (448D)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen HIV-1 infection

Ab Type gp120 CD4BS

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References Gorny & Zolla-Pazner 2004; Nyambi et al. 2000; Wyatt et al. 1998; Li et al. 1997; Manca et al. 1995a; Forthal et al. 1995; Laal et al. 1994; Spear et al. 1993; McKeating et al.

1992c; Karwowska et al. 1992a

Keywords ADCC, antibody binding site definition and exposure, antibody interactions, complement, enhancing activity, review, structure, subtype comparisons, variant cross-recognition or cross-neutralization

- 448-D: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (**review**)
- 448-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 CD4BS MAbs - CD4BS MAbs bound consistently to most isolates of clade D, but poorly to isolates of other clades with the exception of broadly reactive MAb IgG1b12. Nyambi et al. [2000] (variant crossrecognition or cross-neutralization, subtype comparisons)
- 448-D: Summary of the implications of the crystal structure of the core of gp120 bound to CD4 and 17b with what is known about mutations that reduce NAb binding - probable mechanism of neutralization by CD4BS Ab is direct interference with CD4 binding. Wyatt et al. [1998] (structure)
- 448-D: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env. Li et al. [1997] (variant cross-recognition or cross-neutralization)
- 448-D: Neutralizing activity, positive ADCC activity, and no viral enhancing activity. Forthal et al. [1995] (ADCC, enhancing activity)
- 448-D: Virions complexed to gp120 Ab facilitate presentation of p66 RT epitopes to Th cells. Manca et al. [1995a]
- 448-D: Dissociation constant gp120 IIIB 0.029 neutralizes IIIB, acts synergistically with anti-V3 MAb 447-52D. Laal et al. [1994] (antibody interactions)
- 448-D: Did not mediate deposition of complement component C3 on HIV infected cells. Spear et al. [1993] (complement)
- 448-D: Conformational reactive with IIIB gp120 in RIP, but not WB assay. Karwowska et al. [1992a] (antibody binding site definition and exposure)
- 448-D: Called 448D blocks gp120-CD4 binding substitutions at gp120 residues 88, 113, 117, 257, 368 and 370 reduce binding - epitope similar to rat MAbs 39.13g and 39.3b. McKeating et al. [1992c] (antibody binding site definition and exposure)

**No.** 1044

MAb ID 46D2/D5

**HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** 

Subtype B

Neutralizing no

Immunogen vaccine

> HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

**Ab Type** gp120 CD4BS

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References He et al. 2002

• 44D2/D5: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human  $IgG2\kappa$ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—6 anti-CD4BS MAbs competed with anti-CD4BS MAb 5145A, blocked sCD4 binding and were conformationally sensitive—4/6 could neutralize the autologous strain SF162, and were broadly cross-reactive with B clade R5 and X4 strains (not E clade)—44D2/D5 could not neutralize autologous SF162, and while it was cross-reactive, it was at lower affinity. He et al. [2002]

No. 1045

**MAb ID** 48-16

**HXB2 Location** Env Author Location gp120

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection **Species (Isotype)** human ( $IgG\kappa$ ) Ab Type gp120 CD4BS

1995

Keywords antibody binding site definition and exposure, binding affinity, review, variant crossrecognition or cross-neutralization

- 48-16: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database. Most neutralize TCLA strains only, 48-16 is one of four that are non-neutralizing. Gorny & Zolla-Pazner [2004] (review)
- 48-16: Broadly cross-reactive, reacts outside the CD4 binding site and V3 region—competes with sera from 45 seropositive subjects—binding affinity  $2-5 \times 10^{-9}$  M. Fevrier *et al.* [1995] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, binding affinity)

**No.** 1046

**MAb ID** 50-61A

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection **Species (Isotype)** human ( $IgG\kappa$ ) Ab Type gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Fevrier et al.

1995

Keywords binding affinity, review, variant crossrecognition or cross-neutralization

Vector/Type: protein Strain: B clade SF162 • 50-61A: Neutralizes lab strains LAI and SF2 – competes with sera from 45 seropositive subjects – binding affinity 2.4 x 10<sup>-10</sup> M. (variant cross-recognition or cross-neutralization, binding affinity)

> • 50-61A: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)

> > **No.** 1047

**MAb ID** 5145A

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Subtype B

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 CD4BS

Research Contact Abraham Pinter, Public Health Research In-

stitute, Newark, NJ, 07103. pinter@phri.org.

References Pinter et al. 2005; Pinter et al. 2004; Gorny & Zolla-Pazner 2004; He et al. 2002; Alsmadi & Tilley 1998; Pincus et al. 1996; Warrier et al.

1996; Pinter et al. 1993a

Keywords ADCC, antibody binding site definition and exposure, antibody interactions, immunotoxin, variant cross-recognition or cross-

neutralization

- References Gorny & Zolla-Pazner 2004; Fevrier et al. 5145A: This study is about the MAb C108g, and 5145A was a control. 5145A is a disulfide-dependent epitope in the CD4 binding domain that is lost after reduction; C108g, contrary to earlier reports, was also shown to require disulfide bonds. Pinter et al. [2005] (antibody binding site definition and exposure)
  - 5145A: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization)
  - 5145A: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Three anti-CD4BS MAbs were tested, including IgG1b12, which neutralizes both JRFL and SF162. The affinities for IgG1b12 and 5145A were similar for both JRFL and SF612, but 1125A bound with 2.5 fold higher affinity to SF162. 5145A and 1125H both preferentially neutralize SF162, but not JRFL, and the CD4BS is more sensitive to neutralization in the context of the SF162 V1V2 loop. This was also true for neutralization by sCD4. Pinter et al. [2004] (variant cross-recognition or cross-neutralization)
  - 5145A: Transgenic mice carrying human genes allowing production of fully human MAbs were used to rapidly create a

panel of anti-HIV gp120 MAb producing hybridomas by immunization with HIV SF162 gp120 - the previously described human MAbs 5145A, 4117C and 697D were used as controls. **Species (Isotype)** human (IgG1 $\kappa$ ) He et al. [2002]

- 5145A: A study of 6 anti-Env MAbs and their ability to bind Research Contact Susan or direct ADCC against target cells infected with IIIB, MN, SF-2, and RF – bound and directed lysis against all four strains. Alsmadi & Tilley [1998] (ADCC)
- 5145A: A panel of immunotoxins were generated by linking Env MAbs to ricin A – immunotoxins mediated cell killing, but killing was not directly proportional to binding. Pincus et al. [1996] (**immunotoxin**)
- 5145A: Synergistic neutralization of HIV-1 when combined with anti-V2 MAb C108G. Warrier et al. [1996] (antibody interactions)
- 5145A: Potent and broadly cross-reactive neutralization of lab strains. Pinter et al. [1993a] (variant cross-recognition or cross-neutralization)

**No.** 1048

**MAb ID** 558-D

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

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References Gorny & Zolla-Pazner 2004; Nyambi et al.

1998; McKeating et al. 1992c

Keywords antibody binding site definition and exposure, review, subtype comparisons, variant cross-

recognition or cross-neutralization

- 558-D: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 558-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H - CD4BS Abs tended to bind weakly without clade specificity to virions, but bound well to soluble gp120 - 558-D did not bind to any B clade viruses, and weakly bound to clade A, C, and G isolates - 559/64-D, 558-D and 1202-D had similar reactivities. Nyambi et al. [1998] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 558-D: Blocks gp120-CD4 binding binds a panel of mutants all except for 256 S/Y and 262 N/T, which are probably conformationally disruptive. McKeating et al. [1992c] (antibody binding site definition and exposure)

No. 1049

**MAb ID** 559/64-D (559, 559-64D)

**HXB2 Location** Env

Author Location gp120 (LAI)

**Epitope** Subtype B **Neutralizing** L

Immunogen HIV-1 infection

Ab Type gp120 CD4BS

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References Gorny & Zolla-Pazner 2004; Zwick et al.

2003; York et al. 2001; Hioe et al. 2001; Nyambi et al. 2000; Hioe et al. 2000; Gorny et al. 2000; Nyambi et al. 1998; Hioe et al. 1997b; Hioe et al. 1997a; Jeffs et al. 1996; Forthal et al. 1995; Stamatatos & Cheng-Mayer 1995; Spear et al. 1993; McKeating

et al. 1992c; Karwowska et al. 1992a

Keywords ADCC, antibody binding site definition and exposure, antibody interactions, assay development, complement, enhancing activity, review, subtype comparisons, variant crossrecognition or cross-neutralization

- 559/64D: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 559/64D: called 559-64D: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick et al. [2003] (antibody interactions)
- 559/64-D: CD4BS MAbs when complexed with gp120, inhibit proliferation of gp120-specific CD4 T-cells and IFNγ production—anti-CD4BS MAbs inhibit gp120 presentation by altering the uptake and/or processing of gp120 by the APCs, not by blocking of gp120 attachment to CD4 on the surface of APCs. Hioe et al. [2001]
- 559/64-D: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4 induced or CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding. York et al. [2001] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 559/64-D: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared - no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer - CD4BS MAbs 559/64-D, 654-D, 729-D, 9CL and 1331E bound with a 5-13 fold preference for the oligomer.

posure, variant cross-recognition or cross-neutralization)

- 559/64-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses – anti-CD4 binding site MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells - CD4BS MAbs 654-D, 559/64-D, 588-D, 830-D, 1027-30-D, and 1202-30D strongly diminished proliferation. Hioe et al. [2000]
- 559/64-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 CD4BS MAbs -CD4BS MAbs bound consistently to most isolates of clade D, but poorly to isolates of other clades with the exception of broadly reactive MAb IgG1b12. Nyambi et al. [2000] (subtype comparisons)
- 559/64-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H - CD4BS Abs tended to bind weakly without clade specificity to virions, but bound well to soluble gp120 – 559/64-D did not bind to any B clade viruses, and weakly bound clade A, C, and G isolates - 559/64- Research Contact Dr. Abraham Pinter, Public Health Research D, 558-D and 1202-D had similar reactivities. Nyambi et al. [1998] (antibody binding site definition and exposure, subtype comparisons)
- 559/64-D: Used in the development of resting cell neutralization assay. Hioe *et al.* [1997a] (**assay development**)
- 559/64-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs – BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) – isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations – US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe et al. [1997b] (variant cross-recognition or cross-neutralization)
- 559/64-D: Called 559 slight, not significant increased binding when V1/V2 or V1/V2 and V3 were deleted from gp120. Jeffs et al. [1996] (antibody binding site definition and exposure)
- 559/64-D: Neutralizing activity, no ADCC activity, and no viral enhancing activity. Forthal et al. [1995] (ADCC, enhancing activity, variant cross-recognition or cross-neutralization)
- 559/64-D: Called 559-64D The binding of conformationdependent anti-V2, anti-V3, and anti-CD4BS MAbs to monomeric and virion-associated gp120 from HIV-1 isolates with differences in cell tropism was studied - CD4BS loop epitopes are somewhat occluded in the oligomeric gp120 epitopes on the virion surface relative to the gp120 monomer as indicated by an increase in the half-maximal binding values to macrophage-tropic isolates SF162 and SF128a and to T-cell tropic SF2 - binding of anti-CD4BS MAbs to SF2 resulted in a significant amount of dissociation of gp120 from virion surface. Stamatatos & Cheng-Mayer [1995] (antibody binding site definition and exposure)

Gorny et al. [2000] (antibody binding site definition and ex- • 559/64-D: Did not mediate deposition of complement component C3 on HIV infected cells. Spear et al. [1993] (complement)

> • 559/64-D: Conformational – reactive with IIIB gp120 in RIP, but not WB assay. Karwowska et al. [1992a] (antibody binding site definition and exposure)

> > **No.** 1050

MAb ID 55D5/F9

**HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** Subtype B Neutralizing L Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 CD4BS

Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002 Keywords review, variant cross-recognition or crossneutralization

- 55D5/F9: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database. Most neutralize TCLA strains only, this is one of four that are non-neutralizing. Gorny & Zolla-Pazner [2004] (review)
- 55D5/F9: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human  $IgG2\kappa$ MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120-6 anti-CD4BS MAbs competed with anti-CD4BS MAb 5145A, blocked sCD4 binding and were conformationally sensitive—4/6 could neutralize the autologous strain SF162, and were broadly cross-reactive with B clade R5 and X4 strains (not E clade)—55D5/F9 was one of these four MAbs. He et al. [2002] (variant cross-recognition or cross-neutralization)

**No.** 1051

MAb ID 588-D (588)

**HXB2 Location** Env Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 CD4BS

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu), Med

Center, NY, NY

References Nyambi et al. 2000; Hioe et al. 2000; Nyambi

et al. 1998; Jeffs et al. 1996; Moore & Ho 1993; Buchbinder et al. 1992; Karwowska

et al. 1992a

- 588-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses anti-CD4 binding site MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells CD4BS MAbs 654-D, 559/64-D, 588-D, 830-D, 1027-30-D, and 1202-30D strongly diminished proliferation. Hioe *et al.* [2000]
- 588-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 CD4BS MAbs CD4BS MAbs bound consistently to most isolates of clade D, but poorly to isolates of other clades with the exception of broadly reactive MAb IgG1b12. Nyambi *et al.* [2000]
- 588-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H CD4-BS Abs tended to bind weakly without clade specificity to virions, but bound well to soluble gp120 588-D did not bind to any B clade viruses, and weakly bound a clade A, C, and G clade isolate 559/64-D, 558-D and 1202-D reacted had similar reactivities. Nyambi *et al.* [1998]
- 588-D: Called 588 slight, not significant increased binding when V1/V2 or V1/V2 and V3 were deleted from gp120. Jeffs et al. [1996]
- 588-D: Weak neutralization of IIIB strong inhibition of HIV+ human sera binding to IIIB gp120. Moore & Ho [1993]
- 588-D: 4-fold increase in neutralization potency for 588-D when combined 1:1 with human MAb 447-D. Buchbinder *et al*. [1992]
- 588-D: Conformational reactive with IIIB gp120 in RIP, but not WB assay. Karwowska et al. [1992a]

**No.** 1052

**MAb ID** 654-D (654-30D, 654/30D, 654-D100, 654.30D, 654)

**HXB2 Location** Env

Author Location gp120 (LAI)

**Epitope** 

Subtype B

Neutralizing L

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG\kappa$ )

Ab Type gp120 CD4BS

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu), NYU Med Center, NY, NY

References Gorny & Zolla-Pazner 2004; Zwick et al. 2003; Gorny et al. 2002; Verrier et al. 2001; Nyambi et al. 2000; Hioe et al. 2001; Hioe et al. 2000; Gorny et al. 2000; Hioe et al. 1999; Stamatatos & Cheng-Mayer 1998; Nyambi et al. 1998; Schonning et al. 1998; Gorny et al. 1998; Hioe et al. 1997b; Gorny et al. 1997; Stamatatos et al. 1997; Li et al. 1997; Stamatatos & Cheng-Mayer 1995; Gorny et al. 1994; Laal et al. 1994;

Karwowska *et al.* 1993 **Keywords** antibody binding site definition and exposure, antibody interactions, binding affinity, enhancing activity, kinetics, review, subtype com-

parisons, variant cross-recognition or crossneutralization

- 654-D: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 654-D: Called 654-30D. scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick et al. [2003] (antibody interactions)
- 654-D: Called 654: Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated the six new MAbs all bind to the tip of the V3 loop and cross-compete with the MAb 447-52D and are conformationally sensitive MAbs showed cross-clade binding to native, intact virions and the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization studies), 654 (anti-CD4BS used as a conformation-sensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control), and MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades) Gorny et al. [2002]
- 654-D: CD4BS MAbs when complexed with gp120, inhibit proliferation of gp120-specific CD4 T-cells and IFN gamma production anti-CD4BS MAbs inhibit gp120 presentation by altering the uptake and/or processing of gp120 by the APCs, not by blocking of gp120 attachment to CD4 on the surface of APCs. Hioe et al. [2001]
- 654-D: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6—six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D, while six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281—no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier *et al.* [2001] (antibody interactions, variant cross-recognition or cross-neutralization)
- 654-D: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer CD4BS MAbs 559/64-D, 654-D, 729-D, 9CL and 1331E bound with a 5-13 fold preference for the oligomer. Gorny et al. [2000] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 654-D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses CD4BS MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells MAb 654-D

strongly diminished proliferation – there is a discrepancy in isotyping this antibody, previous reports indicated IgG1kappa, while Hioe suggests it is IgG1lambda. Hioe *et al.* [2000]

- 654-D: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 CD4BS MAbs CD4BS MAbs bound consistently to most isolates of clade D, but poorly to isolates of other clades with the exception of broadly reactive MAb IgG1b12 654-D had the weakest binding among CD4BS MAbs, binding to only 4/26 isolates. Nyambi *et al.* [2000] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 654-D: The presence of leukocyte function-associated molecule

   (LFA-1) promotes virus infectivity and hinders neutralization, and anti-LFA-1 MAbs can enhance the neutralizing effect of anti-HIV V3 MAb 447-52D and anti-HIV CD4BS MAb IgG1b12 non-neutralizing anti-HIV CD4BS MAb 654-D did not become neutralizing in the presence of anti-LFA-1 MAbs. Hioe *et al.* [1999]
- 654-D: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H CD4-BS Abs tended to bind very weakly without clade specificity to virions, but bound well to soluble gp120 654-D bound only to JRFL. Nyambi *et al.* [1998] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 654-D: Called 654-D100 654-D100 and IgG1b12 neutralized viruses HIV-BRU and a mutated virus that lacks the V3 loop glycan equally effectively in contrast, sera from guinea pigs immunized with BRU gp120 neutralize viruses more effectively that lack the V3 glycan. Schonning *et al.* [1998] (variant cross-recognition or cross-neutralization)
- 654-D: Called 654.30D deleting the V2 loop of neutralization-resistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F deletion of V2 but not V1 slightly allowed neutralization by CD4BS MAb 654.30D. Stamatatos & Cheng-Mayer [1998] (antibody binding site definition and exposure, subtype comparisons)
- 654-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe *et al.* [1997b] (variant cross-recognition or cross-neutralization)
- 654-D: Called 654-30D One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env. Li et al. [1997] (variant cross-recognition or cross-neutralization)

- 654-D: Anti-CD4 BS MAb 654-30D and IgG1b12 have comparable binding affinities, neither mediates gp120-virion dissociation, but IgG1b12 can neutralize SF128A and SF162 and 654-D cannot 654-D actually enhances infection by both viruses in primary macrophages. Stamatatos *et al.* [1997] (**enhancing activity, binding affinity**)
- 654-D: Called 654-30D The binding of conformation-dependent anti-V2, anti-V3, and anti-CD4BS MAbs to monomeric and virion-associated gp120 from HIV-1 isolates with differences in cell tropism was studied CD4BS loop epitopes are somewhat occluded in the oligomeric gp120 epitopes on the virion surface relative to the gp120 monomer as indicated by an increase in the half-maximal binding values to macrophage-tropic isolates SF162 and SF128a and to T-cell tropic SF2 binding of anti-CD4BS MAbs to SF2 resulted in a significant amount of dissociation of gp120 from virion surface. Stamatatos & Cheng-Mayer [1995] (antibody binding site definition and exposure)
- 654-D: Mild oxidation of carbohydrate moieties inhibits binding. Gorny *et al.* [1994] (**antibody binding site definition and exposure**)
- 654-D: Dissociation constant gp120 IIIB 0.008 neutralizes IIIB, acts synergistically with anti-V3 MAb 447-52D reported to be human(IgG1lambda) Laal *et al.* [1994] (antibody interactions, kinetics)

**No.** 1053

MAb ID 67G6/C4

**HXB2 Location** Env

Author Location gp120 (SF162)

Epitope

Subtype B

Neutralizing no Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 CD4BS

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He *et al.* 2002 Keywords review, variant cross-recognition or cross-neutralization

- 67G6/C4: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database.
   Most neutralize TCLA strains only, this MAb is one of four that are non-neutralizing. Gorny & Zolla-Pazner [2004] (review)
- 67G6/C4: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2 $\kappa$  MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120—6 anti-CD4BS MAbs competed with anti-CD4BS MAb 5145A, blocked sCD4 binding and were conformationally sensitive—4/6 could neutralize the autologous strain SF162, and were broadly cross-reactive with B clade R5 and X4 strains (not E clade)—67G6/C4 could not neutralize autologous SF162, and its binding was strain-specific. He *et al.* [2002] (variant cross-recognition or cross-neutralization)

No. 1054

**MAb ID** 729-D (729-30D)

**HXB2 Location** Env

Author Location gp120 (LAI)

**Epitope** 

Subtype B

**Neutralizing** L

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**Ab Type** gp120 CD4BS **Research Contact** Susan

Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu), NYU Med

Center, NY, NY

**References** Gorny & Zolla-Pazner 2004; Gorny *et al.* 2000; Parren *et al.* 1997b; Li *et al.* 1997;

D'Souza et al. 1997; Laal et al. 1994

**Keywords** antibody binding site definition and exposure, antibody interactions, kinetics, review, variant cross-recognition or cross-neutralization

- 729-D: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 729-D: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer CD4BS MAbs 559/64-D, 654-D, 729-D, 9CL and 1331E bound with a 5-13 fold preference for the oligomer. Gorny *et al.* [2000] (antibody binding site definition and exposure)
- 729-D: In a multilaboratory blinded study, failed to consistently neutralize any of nine B clade primary isolates reported here to have a lambda light chain, but originally reported in Laal *et al.* [1994] to be IgG1kappa D'Souza *et al.* [1997]. D'Souza *et al.* [1997]; Laal *et al.* [1994] (variant cross-recognition or cross-neutralization)
- 729-D: Called 720-30D one of 14 human MAbs tested for ability to neutralize chimeric SHIV-vpu+, which expressed HIV-1 IIIB env. Li *et al.* [1997] (variant cross-recognition or cross-neutralization)
- 729-D: Neutralizes TCLA strains, but not primary isolates.
   Parren et al. [1997b] (variant cross-recognition or cross-neutralization)
- 729-D: Dissociation constant gp120 IIIB 0.025 neutralizes IIIB, acts synergistically with anti-V3 MAb 447-52D. Laal et al. [1994] (antibody interactions, kinetics)

**No.** 1055

**MAb ID** 830D (830-D)

**HXB2 Location** Env

**Author Location** gp120

Epitope

Neutralizing L

**Immunogen** 

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 CD4BS

**References** Gorny & Zolla-Pazner 2004; Hioe *et al.* 2000; Wyatt *et al.* 1998; Hioe *et al.* 1997b

**Keywords** review, structure, variant cross-recognition or cross-neutralization

- 830D: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 830D: Ab responses, because of their capacity to alter antigen uptake and processing, can influence helper T cell responses anti-CD4 binding site MAbs or serum Ig from HIV+ individuals inhibited proliferative responses of gp120 specific T cells CD4BS MAbs 654-D, 559/64-D, 588-D, 830-D, 1027-30-D, and 1202-30D strongly diminished proliferation. Hioe *et al.* [2000]
- 830D: Summary of the implications of the crystal structure of the core of gp120 bound to CD4 and 17b with what is known about mutations that reduce NAb binding – probable mechanism of neutralization by CD4BS Ab is direct interference with CD4 binding. Wyatt et al. [1998] (structure)
- 830D: Called 830-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe *et al.* [1997b] (variant cross-recognition or cross-neutralization)

No. 1056

MAb ID 9CL

**HXB2 Location** Env

Author Location gp120 (LAI)

Epitope

Subtype B

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu), NYU Med

Center, NY, NY

References Gorny & Zolla-Pazner 2004; Gorny et al.

2000

**Keywords** antibody binding site definition and exposure, review

- 9CL: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- 9CL: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer CD4BS MAbs 559/64-D, 654-D, 729-D, 9CL and 1331E bound with a 5-13 fold preference for the oligomer.

Gorny et al. [2000] (antibody binding site definition and exposure)

**No.** 1057

MAb ID BM12

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

References Kessler et al. 1995

• BM12: Broad cross-clade neutralization of primary isolates – additive effect in combination with MAb 2F5. Kessler *et al.* [1995]

No. 1058

MAb ID D20

HXB2 Location Env

Author Location gp120 (IIIB)

Epitope Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

 $\boldsymbol{Species}\;(\boldsymbol{Isotype})\;\;\text{mouse}\;(\boldsymbol{IgG})$ 

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and In-

fectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1997; Otteken et al. 1996; Richardson et al. 1996; Broder

et al. 1994; Earl et al. 1994

Keywords antibody binding site definition and exposure,

antibody generation

- D20: A comparison of 25 gp120 specific, conformation dependent MAbs was done D20 is part of a group of MAbs labeled A1 all A1 MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, blocked CD4 binding, were sensitive to mutations in gp120 positions 368 and 370 that directly contact CD4. Sugiura *et al.* [1999] (antibody binding site definition and exposure)
- D20: Used for comparison in a study of gp41 antibodies D20 binds to a greater extent to cell surface expressed Env than any of 38 conformation dependent anti-gp41 MAbs. Earl *et al.* [1997] (antibody binding site definition and exposure)
- D20: Pulse label experiments of 4 MAbs (D20, D27, T20, and T22) binding to noncleavable gp160 revealed that these anti-CD4 MAbs bound with a delay, and that the epitope formed with a t 1/2 of about 10 minutes. Otteken *et al.* [1996]
- D20: Human sera blocked binding in oligomeric ELISA assay to a similar extent for gp41 MAbs D20, D43, D61, and T4. Richardson *et al.* [1996]
- D20: Binding completely blocked by pooled human sera. Broder *et al.* [1994]
- D20: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994] (**antibody generation**)

**No.** 1059

MAb ID D21

HXB2 Location Env

**Author Location** gp120 (IIIB)

Epitope

Neutralizing

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and In-

fectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- D21: A comparison of 25 gp120 specific, conformation dependent MAbs was done D21 is part of a group of MAbs labeled A1 all A1 MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, blocked CD4 binding, were sensitive to mutations in gp120 positions 368 and 370 that directly contact CD4. Sugiura *et al.* [1999]
- D21: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

**No.** 1060

MAb ID D24

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and In-

fectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- D24: A comparison of 25 gp120 specific, conformation dependent MAbs was done D24 is part of a group of MAbs labeled B-I, that had limited cross-reactivity of seven clade B isolates BH8 and B-al were they only two that consistently reacted with B-I MAbs B-I MAbs fully blocked CD4 binding. Sugiura et al. [1999]
- D24: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

**No.** 1061

MAb ID D25

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope

Neutralizing

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

**Ab Type** gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and In-

fectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

D25: A comparison of 25 gp120 specific, conformation dependent MAbs was done – D25 is part of a group of MAbs labeled A1 – all A1 MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, blocked CD4 binding, were sensitive to mutations in gp120 positions 368 and 370 that directly contact CD4. Sugiura *et al.* [1999]

D25: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1062

MAb ID D28

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope Neutralizing no Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- D28: A comparison of 25 gp120 specific, conformation dependent MAbs was done D28 is part of a group of MAbs labeled B-I, that had limited cross-reactivity of seven clade B isolates BH8 and B-al were they only two that consistently reacted with B-I MAbs B-I MAbs fully blocked CD4 binding. Sugiura et al. [1999]
- D28: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1063

MAb ID D35

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- D35: A comparison of 25 gp120 specific, conformation dependent MAbs was done D35 is part of a group of MAbs labeled B-I, that had limited cross-reactivity of seven clade B isolates BH8 and B-al were they only two that consistently reacted with B-I MAbs B-I MAbs fully blocked CD4 binding. Sugiura et al. [1999]
- D35: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

No. 1064

MAb ID D39

HXB2 Location Env

Author Location gp120 (IIIB)

**Epitope Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- D39: A comparison of 25 gp120 specific, conformation dependent MAbs was done D39 is part of a group of MAbs labeled A1 all A1 MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, blocked CD4 binding, were sensitive to mutations in gp120 positions 368 and 370 that directly contact CD4. Sugiura *et al.* [1999]
- D39: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1065

MAb ID D42

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and In-

fectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- D42: A comparison of 25 gp120 specific, conformation dependent MAbs was done D42 is part of a group of MAbs labeled B-I, that had limited cross-reactivity of seven clade B isolates BH8 and B-al were they only two that consistently reacted with B-I MAbs B-I MAbs fully blocked CD4 binding. Sugiura *et al.* [1999]
- D42: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

**No.** 1066

MAb ID D52

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope

Neutralizing

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and In-

fectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

D52: A comparison of 25 gp120 specific, conformation dependent MAbs was done – D52 is part of a group of MAbs labeled B-I, that had limited cross-reactivity – of seven clade B isolates BH8 and B-al were they only two that consistently reacted with B-I MAbs – B-I MAbs fully blocked CD4 binding. Sugiura et al. [1999]

• D52: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

**No.** 1067

MAb ID D53

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope Neutralizing

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- D53: A comparison of 25 gp120 specific, conformation dependent MAbs was done D53 is part of a group of MAbs labeled B-I, that had limited cross-reactivity of seven clade B isolates BH8 and B-al were they only two that consistently reacted with B-I MAbs B-I MAbs fully blocked CD4 binding. Sugiura et al. [1999]
- D53: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1068

MAb ID D60

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope

Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

**References** Sugiura *et al.* 1999; Richardson *et al.* 1996; Earl *et al.* 1994

- D60: A comparison of 25 gp120 specific, conformation dependent MAbs was done D60 is part of a group of MAbs labeled B-I, that had limited cross-reactivity of seven clade B isolates BH8 and B-al were they only two that consistently reacted with B-I MAbs B-I MAbs fully blocked CD4 binding. Sugiura et al. [1999]
- D60: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1069

MAb ID DA48

**HXB2 Location** Env

Author Location gp120 (BRU)

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

**Ab Type** gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Sullivan et al.

1998a; Parren et al. 1998a

**Keywords** antibody binding site definition and exposure, antibody generation, binding affinity, review, variant cross-recognition or cross-neutralization

- DO8i: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- DA48: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13 > DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren *et al.* [1998a] (**binding affinity**)
- DA48: The HIV-1 virus YU2 entry can be enhanced by MAbs binding to the CD4BS, V3 loop, and CD4i epitopes the activation for this enhanced entry state could be conferred on HxB2 by introducing the YU2 V3 loop, or the YU2 V3 and V1/V2 loops a similar effect is observed by sub-neutralizing concentrations of sCD4 and the effect is dependent of CCR5 Fab Ab fragment DA48 also enhances YU2 entry, ruling out Fc interactions or Env cross-linking as a mechanism while DA48 enhances YU2, it neutralizes HXBc2 DA48 was obtained by panning libraries derived from bone marrow from a >15 year long term non-progressor against BRU gp120. Sullivan et al. [1998a] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization)

**No.** 1070

MAb ID DO8i

**HXB2 Location** Env

Author Location gp120 (BRU)

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

References Sullivan et al. 1998a; Parren et al. 1998a

• DO8i: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13 > DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization

were correlated for both Fabs and MAbs - authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a]

• DO8i – the HIV-1 virus YU2 entry can be enhanced by MAbs binding to the CD4BS, V3 loop, and CD4i epitopes - the activation for this enhanced entry state could be conferred on HxB2 by introducing the YU2 V3 loop, or the YU2 V3 and V1/V2 loops – a similar effect is observed by sub-neutralizing concentrations of sCD4 and the effect is dependent of CCR5 - Fab fragment DO8i also enhances YU2 entry, ruling out Fc interactions or Env cross-linking as a mechanism, while neutralizing HXBc2 – DO8i was obtained by panning libraries derived from bone marrow from a long term asymptomatic donor against BRU gp120. Sullivan et al. [1998a]

**No.** 1071 **MAb ID** F105 (F-105) **HXB2 Location** Env Author Location gp120 **Epitope** Neutralizing L Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**Ab Type** gp120 CD4BS

Research Contact Marshall Posner, Boston MA

References Wilkinson et al. 2005; Selvarajah et al. 2005;

Martín-García et al. 2005; Pantophlet et al. 2004; Ling et al. 2004; Biorn et al. 2004; Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Ohagen et al. 2003; Raja et al. 2003; Xiang et al. 2003; Poignard et al. 2003; Pantophlet et al. 2003a; Kwong et al. 2002; Ferrantelli et al. 2004a; Cavacini et al. 2002; Ling et al. 2002; Liu et al. 2002; Ferrantelli & Ruprecht 2002; Zhang et al. 2002; Basmaciogullari et al. 2002; Grundner et al. 2002; Edwards et al. 2002; Xiang et al. 2002b; Chakrabarti et al. 2002; Xu et al. 2002; Yang et al. 2002; York et al. 2001; Kolchinsky et al. 2001; Si et al. 2001; Yang et al. 2000; Park et al. 2000; Fortin et al. 2000; Baba et al. 2000; Robert-Guroff 2000; Oscherwitz et al. 1999a; Cavacini et al. 1999; Giraud et al. 1999; Sugiura et al. 1999; Kropelin et al. 1998; Sullivan et al. 1998a; Brand et al. 1998; Cavacini et al. 1998a; Li et al. 1998; Cavacini et al. 1998b; Wyatt et al. 1998; Wyatt et al. 1997; Cao et al. 1997b; Li et al. 1997; D'Souza et al. 1997; Parren et al. 1997b; Chen et al. 1996; Litwin et al. 1996; Pincus et al. 1996; Wisnewski et al. 1996; McDougal et al. 1996; Wolfe et al. 1996; Jagodzinski et al. 1996; Khouri et al. 1995; Sullivan et al. 1995; Cavacini et al. 1995; Posner et al. 1995; Turbica et al. 1995; Chen et al. 1994a; Earl et al. 1994; Cavacini et al. 1994a; Cavacini et al. 1994b; Cook et al. 1994; Thali et al. 1994; Bagley et al. 1994; Marasco et al. 1993; Watkins et al. 1993; Pincus et al. 1993; Klasse

et al. 1993a; Potts et al. 1993; Montefiori et al. 1993; Wyatt et al. 1993; Cavacini et al. 1993b; Cavacini et al. 1993a; Posner et al. 1993; Moore & Ho 1993; Posner et al. 1992a; Posner et al. 1992b; Wyatt et al. 1992; Marasco et al. 1992; Thali et al. 1992a; Thali et al. 1991; Posner et al. 1991

Keywords ADCC, antibody binding site definition and exposure, antibody generation, antibody interactions, antibody sequence, variable domain, assay development, brain/CSF, co-receptor, complement, enhancing activity, escape, immunoprophylaxis, immunotherapy, immunotoxin, kinetics, mother-to-infant transmission, mucosal immunity, rate of progression, review, structure, subtype comparisons, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or crossneutralization

- F105: No neutralization of primary isolates observed (John Moore, pers comm). (variant cross-recognition or crossneutralization)
- F105: NIH AIDS Research and Reference Reagent Program:
- F105: The HIV-1 Bori-15 variant was adapted from the Bori isolate for replication microglial cells. Bori-15 had increased replication in microglial cells and a robust syncytium-forming phenotype, ability to use low levels of CD4 for infection, and increased sensitivity to neutralization by sCD4 and 17b. Four amino acid changes in gp120 V1-V2 were responsible for this change. Protein functionality and integrity of soluble, monomeric gp120-molecules derived from parental HIV-1 Bori and microglia-adapted HIV-1 Bori-15 was assessed in ELISA binding assays using CD4BS MAbs F105 and IgG1b12, glycanspecific 2G12, and V3-specific 447-52D, and were unchanged. Association rates of sCD4 and 17b were not changed, but dissociation rates were 3-fold slower for sCD4 and 14-fold slower for 17b. Martín-García et al. [2005] (antibody binding site definition and exposure)
- F105: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. CD4BS MAbs except Fab b12 (b6, b3, F105) did not bind to either GDMR or mCHO. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)
- F105: The crystal structure of the Fab fragment from F105 was solved. It has an extended CDR H3 loop, with a Phe at the apex that may recognize the binding pocket of gp120 used by the Phe-42 residue of CD4. The potent NAB IgG1b12 recognizes an overlapping binding site, the main difference is that F105 extends across the interface of the inner and outer domains of gp120 while b12 does not. IgG1b12 also has undergone extensive affinity maturation (45 mutations) while F105 has not (13 mutations) – an average for gp120 MAbs is 22 mutations.

Wilkinson *et al.* [2005] (antibody sequence, variable domain, structure)

- F105: The peptide 12p1 (RINNIPWSEAMM) inhibits direct binding of YU2 gp120 or Env trimer to CD4, CCR5 and MAb 17b in a concentration-dependent allosteric manner. 12p1 is thought to bind to unbound gp120 near the CD4 binding site, with a 1:1 stoichiometry. 12p1 also inhibited MAb F105 binding. presumably because F105 favors an unactivated conformation, but not MAbs 2G12 or b12. The 1:1 stoichiomtry, the fact that the peptide binding site is accessible on the trimer, the non-CD4 like aspect of the binding, and an ability to inhibit viral infection in cell cultures make 12p1 a promising lead for therapeutic design. Biorn *et al.* [2004]
- F105: NAbs against HIV-1 M group isolates were tested for their ability to neutralize 6 randomly selected HIV-1 O group strains. F105 was not particularly effective at neutarlizing HIV-1 group O strains. Ferrantelli *et al.* [2004a] (variant cross-recognition or cross-neutralization)
- F105: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- F105: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of CD4BS MAb F105 was decreased by trypsin, but increased by thrombin. Thrombin may increase HIV-induced cell fusion in blood by causing a conformational activating shift in gp120. Ling *et al.* [2004] (antibody binding site definition and exposure)
- F105: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including F105. Pantophlet *et al.* [2004] (vaccine antigen design)
- F105: Env genes derived from uncultured brain biopsy samples from four HIV-1 infected patients with late-stage AIDS were compared to env genes from PBMC samples. Brain isolates did not differ in the total number or positions of N-glycosylation sites, patterns of coreceptor usage, or ability to be recognized by gp160 and gp41 MAbs. F105 recognized most variants, some from each of the four individuals by gp120 immunoprecipitation. Ohagen *et al.* [2003] (brain/CSF, variant cross-recognition or cross-neutralization)
- F105: Alanine scanning mutagenesis was used to compare substitutions that affected anti-CD4BS NAb b12 rec gp120s were engineered to contain combinations of Alanine substitutions that enhanced b12 binding, and while binding of b12 to these gp120 monomers was generally maintained or increased, binding by five non-neutralizing anti-CD4bs MAbs (b3, b6, F105, 15e, and F91) was reduced or completely abolished. Pan-

tophlet *et al.* [2003a] (antibody binding site definition and exposure)

- F105: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of non-neutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet *et al.* [2003b] (vaccine antigen design)
- F105: Virion capture assays are not a good preditor of neutralization, and the presentation of epitopes using this assay seems to be different from that of functional Envelope spikes on primary isolates F105 and b6 could efficiently block the b12-mediated capture of infectious virions in a virus capture, but did not inhibit b12 neutralization while b12 was potent at neutralizing the three primary virions JR-CSF, ADA, and 89.6, the Abs F105, 19b, and Fab b6 were overall very poor neutralizers. Poignard *et al.* [2003] (antibody interactions)
- F105: This paper shows that binding of CD4BS MAbs to Env blocks the conformational shift that allows co-receptor CCR5 binding and CD4-independent mediated cell fusion. CD4BS MAbs F105, 15e, and IgG1b12 as well as their Fab fragments inhibited CD4-independent binding of the V1/V2 loop-deleted gp120 glycoproteins of R5 HIV-1 isolates ADA, YU2 and JR-FL and to CCR5 in a concentration dependent manner. CD4BS MAbs IgG1b12, F91 and F105 and their Fab counterparts (except for C11, used as a negative control) inhibited CD4-independent JR-FL and YU-2 gp120-CCR5 binding to CCR5-expressing Cf2Th cells and syncytium formation. Raja et al. [2003] (antibody binding site definition and exposure, co-receptor)
- F105: 17b: This paper describes the generation of CD4i MAb E51, that like CD4i MAb 17b, blocks CCR5 binding to sCD4-bound gp120. The substitutions E381R, F383S, R419D I420R, K421D, Q422L, I423S, and Y435S (HXB2 numbering) all severely reduce 17b and E51 binding. All but I423S also diminish CCR5 binding by more than 50%. The mutation F383S also inhibits sCD4 binding and F105 binding, and K421D inhibits F105 binding, but not sCD4. Xiang *et al.* [2003] (antibody binding site definition and exposure)
- F105: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick et al. [2003] (antibody interactions)
- F105: gp120 mutants were used to define the CXCR4 binding site using CXCR4 displayed on paramagnetic proteoliposomes (PMPLs) to reduce non-specific gp120 binding—basic residues in the V3 loop and the β19 strand (RIKQ, positions 419-422) were involved, and deletion of the V1-V2 loops allowed CD4-independent CXCR4 binding—MAbs 17b (CD4i)

and F105 (CD4BS) were used to study conformational changes in the mutants—the affinity of  $\Delta V1$  and  $\Delta V1$ -V2 mutants for F105 was comparable to the wildtype—V3 mutants did not affect F105 binding—the K421A mutation in the  $\beta$ 19 strand dramatically reduced F105 affinity, consistent with what is known about the F105 epitope. Basmaciogullari *et al.* [2002] (antibody binding site definition and exposure)

- F105: This study examined Ab interactions, binding and neutralization with a B clade R5 isolate (92US660) and R5X4 isolate (92HT593). Abs generally bound and neutralized the R5X4 isolate better than the R5 isolate. Anti-gp41 MAb F240 enhanced the binding of CD4BS MAbs IgG1b12 and F105 to both R5X4 and R5 isolates, but had no effect on neutralization. Anti-V3 MAb B4a1 increased CD4BS MAbs IgG1b12 and F105 to R5X4 virions, but only IgG1b12 binding was increased by B4a1 to the R5 isolate, and neutralization was not impacted. Cavacini *et al.* [2002] (**co-receptor**)
- F105: A modified gp140 (gp140deltaCFI), with C-term mutations intended to mimic a fusion intermediate and stabilize trimer formation, retained antigenic conformational determinants as defined by binding to CD4 and to MAbs 2F5, 2G12, F105, and b12, and enhanced humoral immunity without diminishing the CTL response in mice injected with a DNA vaccine. Chakrabarti et al. [2002] (vaccine antigen design)
- F105: Review of NAbs that notes that F105 binds the CD4BS, in combination with other MAbs it can protect some macaques against SHIV infection, and that it has strong ADCC activity.
   Ferrantelli & Ruprecht [2002] (ADCC, antibody interactions, immunoprophylaxis, review)
- F105: HIV-1 gp160δCT (cytoplasmic tail-deleted) proteoliposomes (PLs) containing native, trimeric envelope glycoproteins from R5 strains YU2 and JRFL, and X4 strain HXBc2, were made in a physiologic membrane setting as candidate immunogens for HIV vaccines—2F5 bound to gp160δCT with a reconstituted membrane ten-fold better than the same protein on beads, while such an affinity difference was not seen with F105 and 2G12—anti-CD4BS MAbs IgG1b12 and F105, A32 (C1-C4), C11 (C1-C5), and 39F (V3) MAbs bound gp160δCT PLs indistinguishably from gp160δCT expressed on the cell surface. Grundner *et al.* [2002] (antibody binding site definition and exposure)
- F105: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar. Enthalpy and entropy changes were divergent, but compensated. Not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs (17b, 48d, 1.5e, b6, F105 and F91) had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, but the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. The high values suggest surface burial or protein folding and ordering of amino acids. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking,

- requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong *et al.* [2002] (**antibody binding site definition and exposure**)
- F105: A pseudotyping assay showed that an X4 V3 loop peptide could enhance infectivity of X4 virus, R5 and R5X4 V3 loops peptides could enhance infectivity of an R5 virus, and R5X4 peptides could enhance infectivity of an R5X4 virus. Neither R5 nor R5X4 peptides influenced binding of CD4BS MAbs F105 and Ig1Gb12, but did increase binding of CD4i MAb 17b. Ling et al. [2002] (antibody binding site definition and exposure, co-receptor)
- F105: Review of NAbs that discusses mechanisms of neutralization, passive transfer of NAbs and protection in animal studies, and vaccine strategies. Liu *et al.* [2002] (immunoprophylaxis)
- F105: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations 375 S/W seems to favor a conformation of gp120 closer to the CD4-bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang et al. [2002b] (antibody binding site definition and exposure)
- F105: Passive immunization of neonate macaques with a combination of F105+2G12+2F5 conferred complete protection against oral challenge with SHIV-vpu+ or the combination b12+2G12+2F5 conferred partial protection against SHIV89.6 such combinations may be useful for prophylaxis at birth and against milk born transmission the synergistic combination of IgG1b12, 2G12, 2F5, and 4E10 neutralized a collection of HIV clade C primary isolates. Xu et al. [2002] (immunoprophylaxis, mother-to-infant transmission)
- F105: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin—stabilized oligomer gp140 $\delta$ 683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and C11, A32, and 30D which did not bind the stabilized oligomer. Yang *et al.* [2002] (vaccine antigen design)
- F105: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-

primary strain. Zhang *et al.* [2002] (antibody binding site definition and exposure)

- F105: Mutations in two glcosylation sites in the V2 region of HIV-1 ADA at positions 190 and 197 (187 DNTSYRLINCNTS 199) cause the virus to become CD4-independent and able to enter cells through CCR5 alone – these same mutations tended to increase the neutralization sensitivity of the virus, including to F105. Kolchinsky *et al.* [2001] (antibody binding site definition and exposure)
- F105: SHIV-HXBc2 is a neutralization sensitive non-pathogenic virus, and several *in vivo* passages through monkeys yielded highly pathogenic SHIV KU-1—HXBc2 and the KU-1 clone HXBc2P3.2 differ in 12 amino acids in gp160—substitutions in both gp120 and gp41 reduced the ability of sCD4, IgG1b12, F105 and AG1121 to Env achieve saturation and full occupancy, and neutralize KU-1—17b and 2F5 also bound less efficiently to HXBc2P3.2, although 2G12 was able to bind both comparably. Si *et al.* [2001] (antibody binding site definition and exposure)
- F105: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding. York et al. [2001] (antibody binding site definition and exposure)
- F105: A triple combination of 2F5, F105 and 2G12 effectively neutralized perinatal infection of macaque infants when challenged with SHIV-vpu+ the plasma half-life was 7.2 +/- 2.2 days. Baba *et al.* [2000] (immunoprophylaxis, mother-to-infant transmission)
- F105: Host encoded intercellular adhesion molecule (ICAM-1) is incorporated by the HIV-1 virion and enhances viral infectivity ICAM-1 does not modify virus sensitivity to antibodies 0.5beta or 4.8D or sCD4, but neutralizing ability of F105 was diminished in ICAM bearing virions in the presence of lymphocyte function-association antigen-1 (LFA-1) Ab. Fortin *et al.* [2000]
- F105: Six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form, although F105 was an exception and cannot neutralize either form of MN the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park *et al.* [2000]
- F105: A mini-review of observations of passive administration of IgG NAbs conferring protection against intervenous or vaginal SHIV challenge, that considers why IgG MAbs might protect against mucosal challenge. Robert-Guroff [2000] (immunoprophylaxis, mucosal immunity, review)
- F105: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120

- or gp140 non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang *et al.* [2000] (vaccine antigen design)
- F105: A comparison of 25 gp120 specific, conformation dependent MAbs was done and F105 was used for competition studies F105 did cross-compete with multiple CD4BS specific MAbs, however most could not neutralize even the autologous NL4-3 strains. Sugiura et al. [1999] (antibody interactions)
- F105: Immunoprecipitation of gp120 and gp160 expressed from a rec Semliki Forest virus by F105 and IgG1b12 indicated that the SFV expressed HIV-1 Env was folded appropriately and SVF-HIV-1 Env vaccine gave the strongest anti-HIV-1 Env response in mice, when compared to an HIV-1 Env DNA vaccine and a rgp160 protein. Brand *et al.* [1998] (vaccine antigen design)
- F105: Phase I dose escalation study, single dose of 100 or 500 mg/m2 was given to 4 HIV+ patients sustained levels, no immune response against F105, no toxicity, infused Ab retained function there was no evidence of anti-HIV-1 activity and virus was not diminished at day 1 or 7, by culture or plasma RNA. Cavacini *et al.* [1998b] (kinetics, immunotherapy)
- F105: The MAb F240 binds to the immunodominant region of gp41 and enhances infection in the presence of complement reactivity of F240 is enhanced by preincubation of cells with sCD4 or anti-CD4BS MAb F105. Cavacini *et al.* [1998a] (antibody interactions)
- F105: Anti-C1 region MAb 87-135/9 blocks gp120 interaction with CD4+ cells blocking activity is additive when combined with antibodies which bind in the C4 region of gp120 (F105, 388/389, and b12). Kropelin *et al.* [1998] (antibody interactions)
- F105: Neutralization synergy was observed when the MAbs 694/98-D (V3), 2F5 (gp41), and 2G12 (gp120 discontinuous) were used in combination, and even greater neutralizing potential was seen with the addition of a fourth MAb, F105 (CD4 BS). Li *et al.* [1998] (antibody interactions)
- F105: F105 enhances viral entry of viruses carrying the YU2 envelope glycoproteins, but neutralizes HXBc2. Sullivan *et al.* [1998a] (enhancing activity)
- F105: Summary of the implications of the crystal structure of the core of gp120 bound to CD4 and 17b with what is known about mutations that reduce NAb binding probable mechanism of neutralization by CD4BS Ab is direct interference with CD4 binding. Wyatt *et al.* [1998] (antibody binding site definition and exposure, structure)
- F105: Virus with the V1-V2 loop deleted was viable and more susceptible to neutralization by CD4i MAb 17b, and anti-V3 MAbs 1121, 9284, and 110.4, but not to a CD4BS MAb, F105 or sCD4. Cao *et al.* [1997b] (antibody binding site definition and exposure)
- F105: In a multilaboratory blinded study, failed to neutralize any of nine B clade primary isolates. D'Souza *et al.* [1997] (variant cross-recognition or cross-neutralization)
- F105: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env F105

could only achieve 50% neutralization alone – all Ab combinations tested showed synergistic neutralization – F105 has synergistic response with MAbs 694/98-D (anti-V3), 48d, 2F5, and 2G12, and also with HIVIG. Li *et al.* [1997] (antibody interactions, variant cross-recognition or cross-neutralization)

- F105: Neutralizes TCLA strains, but not primary isolates.
   Parren et al. [1997b] (variant cross-recognition or cross-neutralization)
- F105: Binds both gp120 and soluble gp120+gp41 complex efficiently, suggesting its gp120 epitope is not blocked by gp41 binding does not bind to HXBc2 gp120 if the 19 C-term amino acids, in conjunction with C1 positions 31-93, are deleted. Wyatt *et al.* [1997] (antibody binding site definition and exposure)
- F105: Intracellular co-expression of heavy and light chains of
  the Fab105 fragment MAb F105 was enhanced by inclusion of
  an internal ribosome entry site (IRES) sequence the Fab105
  IRES expression cassette was cloned into an adeno-associated
  virus (AAV) shuttle vector, and transduced into human lymphocytes which were able to produce and secrete the Fab105
  fragments while maintaining normal growth several primary
  HIV-1 patient isolates were effectively blocked. Chen et al.
  [1996] (immunotherapy)
- F105: The sulfated polysaccharide curdlan sulfate (CRDS) binds to the Envelope of T-tropic viruses and neutralizes virus

   deletion of the V3 loop results in less potent inhibition of
   F105 binding by CRDS binding site of F105 described as
   256-257 ST, 368-370 DPE, 421 K, and 470-484 PGGGDMRD-NWRSELY. Jagodzinski *et al.* [1996] (antibody binding site definition and exposure)
- F105: Binding of F105 to oligomeric gp120 occurs despite the fact it cannot neutralize primary isolates. Litwin *et al.* [1996]
- F105: Neutralizes HIV-1 LAI less potently than V3 specific MAbs. McDougal et al. [1996]
- F105: A panel of immunotoxins were generated by linking Env MAbs to ricin A immunotoxins mediated cell killing, but killing was not directly proportional to binding. Pincus *et al.* [1996] (**immunotoxin**)
- F105: F105 is V H4 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996] (antibody sequence, variable domain)
- F105: Phase I study MAb clearance in plasma has a 13 day half-life. Wolfe et al. [1996] (kinetics, immunotherapy)
- F105: Changing heavy chain from IgG1 to IgG3 increased neutralization efficiency. Cavacini *et al.* [1995]
- F105: Biotinylated F105 was used for competition studies with Ab derived from pregnant HIV-1 + women a correlation between maternal anti-CD4 BS Abs overlapping the F105 binding site and lack of HIV-1 transmission to infants was noted. Khouri *et al.* [1995] (mother-to-infant transmission)
- F105: Eight patient phase Ia trial for use as an immunotherapeutic no clinical or biochemical side effects observed, plasma levels of 10 ug/ml maintained for 21 days. Posner *et al.* [1995] (immunotherapy)
- F105: Efficient neutralization of T-cell adapted lines HXBc2 and MN, no neutralization of primary isolates 89.6, ADA and YU2 – even some enhancement of infection of ADA and YU2

was observed. Sullivan *et al.* [1995] (**enhancing activity**, **variant cross-recognition or cross-neutralization**)

- F105: An immunoassay for titrating CD4BS serum antibody was developed using a gp120-coated solid phase and competition with MAb F105 109/110 French HIV-1 + sera and 51/56 HIV-1 + African sera had detectable CD4BS Abs using this assay, demonstrating CD4 binding site conservation among diverse subtypes CD4BS Abs were detected soon after seroconversion and persisted 0/21 HIV-2 + sera reacted, indicating that the HIV-1 and HIV-2 CD4BS Abs are not cross-reactive. Turbica *et al.* [1995] (assay development, subtype comparisons)
- F105: Comparison of MAb F105 sequences with those of MAbs 21h and 15e. Bagley et al. [1994] (antibody sequence, variable domain)
- F105: Administered intravenously to four cynomologus monkeys, plasma pharmacokinetics and biological activity tested. Cavacini *et al.* [1994b] (**kinetics**)
- F105: Fab fragments show reduced capacity to neutralize IIIB, MN, and RF compared to intact IgG1, suggesting bivalent interaction may be important in binding and neutralization. Cavacini *et al.* [1994a] (variant cross-recognition or cross-neutralization)
- F105: A human CD4+ T lymphocyte line was transduced to express Fab fragments of F105 heavy and light chains are joined by an inter-chain linker in the transduced cells infected with HIV-1, the Fab binds intracellularly to the envelope protein and inhibits HIV-1 production secreted Fab fragments neutralize cell-free HIV-1– combined intra- and extracellular binding activities of the expressed Fab make transduced cells resistant to HIV-1 infection and also can protect surrounding lymphocytes by secreting neutralizing antibodies. Chen *et al.* [1994a]; Marasco *et al.* [1993] (variant cross-recognition or cross-neutralization)
- F105: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon anti-CD4 MAbs moderately inhibit gp120 binding to GalCer, possibly through steric hindrance binding of GalCer to gp120 inhibited but did not completely block F105 binding. Cook *et al.* [1994] (brain/CSF)
- F105: Used as a positive control for CD4 BS antibodies in a study of the influence of oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994] (antibody binding site definition and exposure)
- F105: A mutation in gp41, 582 A/T, confers resistance to neutralization (also confers resistance to MAbs 48d, 21h, 15e and 17b). Thali *et al.* [1994] (antibody binding site definition and exposure)
- F105: Additive MN or SF2 neutralization when combined with anti-V3 MAbs 447-52D and 257-D. Cavacini *et al.* [1993a] (antibody interactions)
- F105: Serum from all asymptomatic HIV-1 positive people tested block F105 binding, but only from 27% of symptomatic individuals. Cavacini *et al.* [1993b] (**rate of progression**)
- F105: The gp41 mutation 582(Ala to Thr) results in conformational changes in gp120 that confer neutralization resistance to a class of conformation sensitive neutralizing MAbs required >81 fold higher concentrations to neutralize the mutant than wild type. Klasse *et al.* [1993a] (antibody interactions)

- nated volunteers with V3-loop specific neutralization activity - 2/3 sera demonstrated neutralization synergy, and 3/3 binding/fusion-inhibition synergy. Montefiori et al. [1993] (antibody interactions)
- F105: Called F-105 neutralizes IIIB strong inhibition of HIV+ human sera binding to IIIB gp120. Moore & Ho [1993]
- F105: Ab response in IIIB lab workers was compared to gp160 LAI vaccine recipients – F105 was used as a control – infected lab workers and some of the gp160 vaccinees had a MAb response that could inhibit gp120-CD4 binding, at lower titers than the infected lab workers. Pincus et al. [1993] (vaccinespecific epitope characteristics)
- F105: F105 binds to and neutralizes selected lab strains and 3/9 HIV-1 primary isolates – synergistic enhancement of neutralization by seropositive sera. Posner et al. [1993] (antibody interactions, variant cross-recognition or cross-neutralization)
- · F105: Study of synergy of neutralization and binding comparing F105 and sCD4 with the V3 MAbs: 50.1, 59.1, 83.1, and 58.2 – synergy was observed, and the data suggest that binding of one ligand (F105) can increase the binding of the second (e. g. V3 loop MAbs) due to conformational changes. Potts et al. [1993] (antibody interactions)
- F105: A neutralization escape mutant (HXB2 A281V) was selected by growth of HXB2 in the presence of broadly neutralizing sera – F105 neutralization was not affected by this mutation. Watkins et al. [1993] (escape)
- F105: Binding to Delta V1/2 and Delta V1/2/3 mutant glycoproteins is 2.4- and 13-fold greater, respectively, than binding to wildtype gp120. Wyatt et al. [1993] (antibody binding site definition and exposure)
- F105: MAb cDNA sequence V H4 V71-4 rearranged with a D H D-D fusion product of dlr4 and da4, and with J H5 -V kappa is from the Humvk325 germline gene joined with Jkappa 2. Marasco et al. [1992] (antibody sequence, variable domain)
- F105: F105 mediates ADCC against SF2 through the CD16+ population of PBMC - does not mediate complementdependent cytotoxicity. Posner et al. [1992b] (ADCC, complement)
- F105: Significant enhancement of F105 binding to RF infected cells preincubated with V3-specific MAbs V3-2 and V3-1. Posner et al. [1992a] (antibody interactions)
- F105: Amino acid substitutions that impair F105 neutralization inhibit gp120-CD4 interaction. Thali et al. [1992a] (antibody binding site definition and exposure)
- F105: Precipitation of Delta 297-329 env glycoprotein, which has a deleted V3 loop, is much more efficient than precipitation of wild type. Wyatt et al. [1992] (antibody binding site definition and exposure)
- F105: First description of F105, binds topographically near the CD4-binding site – inhibits binding of free, infectious virions to uninfected HT-H9 cells, but does not react with virus adsorbed to uninfected HT-H9 cells - soluble rCD4 pre-bound to infected cells inhibits F105 binding - F105 inhibits infection of HT-H9 cells in standard neutralization assays with HIV-1 and MN strains. Posner et al. [1991] (antibody binding site definition and exposure, antibody generation)

• F105: Study of synergy between F105 and sera from vacci- • F105: F105 neutralization escape mutants result from changes in amino acids in discontinuous regions: C2, 256-262 and C3, 386-370. Thali et al. [1991] (antibody binding site definition and exposure)

No. 1072

**MAb ID** F91 (F-91)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype)

Ab Type gp120 CD4BS

Research Contact James Robinson, University of Connecticut,

Storrs

References Pantophlet et al. 2004; Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Raja et al. 2003; Pantophlet et al. 2003a; Kwong et al. 2002; Xiang et al. 2002b; Yang et al. 2002; Yang et al. 2000; Fouts et al. 1998; Binley et al. 1998; Parren et al. 1998a; Mondor et al. 1998; Fouts et al. 1997; Moore & Sodroski 1996; Moore et al. 1994b; Moore & Ho 1993

Keywords antibody binding site definition and exposure, antibody interactions, co-receptor, review, subtype comparisons, vaccine antigen design, variant cross-recognition or crossneutralization

- F91: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- F91: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including F91. Pantophlet et al. [2004] (vaccine antigen design)
- F91: Alanine scanning mutagenesis was used to compare substitutions that affected anti-CD4BS NAb b12 - rec gp120s were engineered to contain combinations of Alanine substitutions that enhanced b12 binding, and while binding of b12 to these gp120 monomers was generally maintained or increased, binding by five non-neutralizing anti-CD4bs MAbs (b3, b6, F105, 15e, and F91) was reduced or completely abolished. Pantophlet et al. [2003a] (antibody binding site definition and
- F91: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional Nlinked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and

C5 were also removed, but this compromised b12 binding. Pantophlet *et al.* [2003b] (vaccine antigen design)

- F91: This paper shows that binding of CD4BS MAbs to Env blocks the conformational shift that allows co-receptor CCR5 binding and CD4-independent mediated cell fusion. CD4BS MAbs F105, 15e, and IgG1b12 as well as their Fab fragments inhibited CD4-independent binding of the V1/V2 loop-deleted gp120 glycoproteins of R5 HIV-1 isolates ADA, YU2 and JR-FL and to CCR5 in a concentration dependent manner. CD4BS MAbs IgG1b12, F91 and F105 and their Fab counterparts (except for C11, used as a negative control) inhibited CD4-independent JR-FL and YU-2 gp120-CCR5 binding to CCR5-expressing Cf2Th cells and syncytium formation. Raja *et al.* [2003] (co-receptor)
- F91: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick et al. [2003] (antibody interactions)
- F91:Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar. Enthalpy and entropy changes were divergent, but compensated. Not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs (17b, 48d, 1.5e, b6, F105 and F91) had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, but the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. The high values suggest surface burial or protein folding an ordering of amino acids. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong et al. [2002] (antibody binding site definition and exposure)
- F91: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations 375 S/W seems to favor a conformation of gp120 closer to the CD4-bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang *et al.*

[2002b] (antibody binding site definition and exposure)

- F91: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin stabilized oligomer gp140 delta683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and C11, A32, and 30D which did not bind the stabilized oligomer. Yang *et al.* [2002] (antibody binding site definition and exposure)
- F91: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang *et al.* [2000] (antibody binding site definition and exposure)
- F91: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein (Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer CD4BS MAbs 15e, F91 and IgG1b12 bound better to the deleted protein than to wild type. Binley et al. [1998] (antibody binding site definition and exposure)
- F91: CD4BS MAbs 15e, 21h, and F91 bind with even lower affinity than 205-43-1 and 205-42-15 to JRFL oligomer conclusions of this paper contrast with Parren *et al.* [1998a] Fouts *et al.* [1998]. Fouts *et al.* [1998]; Parren *et al.* [1998a] (antibody binding site definition and exposure)
- F91: Weak inhibition of binding of Hx10 to CD4 positive or negative cells, weakly neutralizing. Mondor et al. [1998]
- F91: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a] (antibody binding site definition and exposure)
- F91: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding F91 bound monomer, did not bind oligomer or neutralize JRFL. Fouts *et al.* [1997] (antibody binding site definition and exposure)
- F91: Unusual pattern of reciprocal enhancement with several anti-V2 and V3 directed MAbs – reciprocal inhibition of other CD4BS MAbs. Moore & Sodroski [1996] (antibody binding site definition and exposure, antibody interactions)
- F91: Has strong cross-reactivity with gp120 monomers from most subtypes, A-F. Moore et al. [1994b] (subtype comparisons)
- F91: Called F-91 neutralizes IIIB reactive with SF-2 gp120 strong inhibition of HIV+ human sera binding to IIIB gp120. Moore & Ho [1993] (variant cross-recognition or cross-neutralization)

**No.** 1073

MAb ID FG39

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

**Ab Type** gp120 CD4BS

References Zwick et al. 2003

Keywords antibody interactions

• FG39: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This is a novel CD4BS Fab first used in this study. Zwick *et al.* [2003] (antibody interactions)

**No.** 1074

MAb ID Fbb14

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

References Zwick et al. 2003

**Keywords** antibody interactions

• Fbb14: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This is a novel CD4BS Fab first used in this study. Fbb14 was unusual among CDBS Abs in that it didn't enhance 4KG5's binding, like b12, but it did not inhibit it either as the other 13 CD4BS Abs did, it remained neutral. Zwick et al. [2003] (antibody interactions)

**No.** 1075

**MAb ID** GP13 (ARP3054)

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Vella et al. 2002; Schutten et al. 1997; Schutten et al. 1996; Wisnewski et al. 1996; Bolmstedt et al. 1996; Schutten et al. 1995b; Schutten et al. 1995a; Bagley et al. 1994; Back et al. 1993; Schutten et al. 1993

**Keywords** antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain, assay development, binding affinity, enhancing activity, escape, review, subtype comparisons, variant cross-recognition or cross-neutralization

- GP13: UK Medical Research council AIDS reagent: ARP3054.
- GP13: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- GP13: Called ARP3054: Herpesvirus saimiri-immortalized CD4+ T lymphocytes (HVS T cells) were used to isolate virus and perform HIV-1 neutralization assays, and compared with a standard PBMC protocol – neutralization sensitivities to a panel of MAbs and to homologous or heterologous plasma/sera were similar for HVS T cells (CN-2 cells) and PBMCs. Vella et al. [2002] (assay development)
- GP13: Neutralized (50%) an SI-env chimeric virus and enhanced (>5 fold) an NSI-env chimeric virus. Schutten *et al.* [1997] (enhancing activity, variant cross-recognition or cross-neutralization)
- GP13: Sera were obtained from guinea pigs vaccinated either with gp160, or with gp160 lacking N-linked glycans at N406, N448, and N463 these sera could block equally well both the CD4 BS MAb GP13 and the V3 MAb F58/H3. Bolmstedt *et al.* [1996] (antibody interactions)
- GP13: IIIB neutralizing MAbs in vitro fail to neutralize in a mouse model in vivo. Schutten et al. [1996]
- GP13: GP13 is V H5 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996] (antibody sequence, variable domain)
- GP13: Neutralizes IIIB only slight inhibition of SI phenotype, and strong enhancement of NSI phenotype chimeric viruses, that incorporated different envs from the same donor. Schutten et al. [1995a] (enhancing activity, variant cross-recognition or cross-neutralization)
- GP13: Neutralizes T-cell adapted viruses but not the SI strain 16.2, despite high binding affinity. Schutten *et al.* [1995b] (variant cross-recognition or cross-neutralization, binding affinity)
- GP13: Mutations in a neutralization resistant isolate obtained by passage of the IIIB isolate in chimpanzees reduced neutralization, but the escape was not as clear as seen with anti-V3 MAbs. Back *et al.* [1993] (escape)
- GP13: Neutralized a broad range of HIV-1 strains from phylogenetically different subfamilies the following gp120 amino acid substitutions strongly inhibit binding: 256(S/Y), 257(T/G), 262(N/T), 368(D/R or K), 370(E/R or Q or D), 384(Y/E). Schutten *et al.* [1993] (antibody binding site definition and exposure, subtype comparisons)

No. 1076

MAb ID GP44

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Wisnewski et al.

1996; Bagley et al. 1994; Schutten et al. 1993

Keywords antibody binding site definition and exposure, antibody sequence, variable domain, review, variant cross-recognition or cross-

neutralization

• GP44: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)

- GP44: GP44 is V H1 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996] (antibody sequence, variable domain)
- GP44: Exhibited a more restricted pattern of neutralizing activity than GP13 and GP68 - the following gp120 amino acid substitutions strongly inhibit binding: 256(S/Y), 257(T/G), 262(N/T), 368(D/R or K), 370(E/R or Q or D) Schutten et al. [1993] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)

No. 1077

MAb ID GP68

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

Ab Type gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Guillon et al.

2002b; Wisnewski et al. 1996; Schutten et al. 1995a; Bagley et al. 1994; Klasse et al. 1993a;

Schutten et al. 1993

Keywords antibody binding site definition and exposure, antibody sequence, variable domain, enhancing activity, review, variant cross-recognition

or cross-neutralization

• GP68: UK Medical Research Council AIDS reagent: ARP3055.

- GP68: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- GP68: The affect of Ab binding on infectivity was studied by pseudotyping three related envs with different phenotypes – R5 viruses were preferentially enhanced, not X4 – the V3 region was the main determinant of Ab-mediated enhancement and modulation of the interaction between CCR5 and gp120 is critical – tests with MAbs anti-V3 391/95-D and CD4BS-specific GP68 indicate that Ab specificity did not determine whether

or not infectivity was enhanced or neutralized, rather the phenotype was determined by Envelope conformation. Guillon et al. [2002b] (enhancing activity, variant cross-recognition or cross-neutralization)

- GP68: GP68 is V H1 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski et al. [1996] (antibody sequence, variable domain)
- GP68: Neutralizes IIIB only slight inhibition of SI phenotype, and strong enhancement of NSI phenotype chimeric viruses, that incorporated different envs from the same donor. Schutten et al. [1995a] (variant cross-recognition or crossneutralization)
- GP68: The gp41 mutation 582(Ala to Thr) results in conformational changes in gp120 that confer neutralization resistance to a class of conformation sensitive neutralizing MAbs - GP68 required markedly higher concentrations to neutralize the mutant than wild type. Klasse et al. [1993a] (antibody binding site definition and exposure)
- GP68: Neutralized a broad range of HIV-1 lab strains from phylogenetically different subfamilies – the following gp120 amino acid substitutions strongly inhibit binding: 117(K/W), 256(S/Y), 257(T/G), 262(N/T), 368(D/R or K), 370(E/R or Q), 384(Y/E), 435(Y/H) Schutten et al. [1993] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)

**No.** 1078

MAb ID HF1.7

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen anti-idiotype

Species (Isotype) mouse (IgM)

Ab Type gp120 CD4BS

References Chanh et al. 1987

• HF1.7: An anti-Id antibody stimulated by anti-CD4 MAb Leu-3a binds to recombinant gp160, suggesting HF1.7 mimics CD4. Chanh et al. [1987]

**No.** 1079

**MAb ID** HT5 (205-43-1)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Subtype B

**Neutralizing** L (weak)

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

Research Contact Ciba-Geigy AG (Basel, Switzerland), and

Tanox Biosystems, Houston, Texas

References Pugach et al. 2004; Gorny & Zolla-Pazner 2004; Herrera et al. 2003; Grovit-Ferbas et al. 2000; Parren et al. 1998a; Fouts et al. 1998; Fouts et al. 1997; Moore et al. 1995a; Moore

et al. 1994b

Keywords reversion, viral fitness, variant crossrecognition or cross-neutralization

- HT5: Also called 205-43-1. This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004]
- HT5: Called 205-43-1: A primary isolate, CC1/85, was passaged 19 times in PBMC and gradually acquired increased sensitivity to FAb b12 and sCD4 that was attributed to changes in the the V1V2 loop region, in particular the loss of a potential glycosylation site. The affinity for sCD4 was unchanged in the monomer, suggesting that the structural impact of the change was manifested at the level of the trimer. The passaged virus, CCcon19, retained an R5 phenotype and its neutralization susceptibility to other Abs was essentially the same as CC1/85. Three CD4BS MAbs, 205-46-9, 205-42-15, and 205-43-1, did not neutralize either the primary or passaged variant. Pugach *et al.* [2004] (reversion, viral fitness, variant cross-recognition or cross-neutralization)
- HT5: Called 205-43-1 CD4BS MAbs b12 (neutralizing) and 205-42-15, 204-43-1, 205-46-9 (nonneutralizing) all cross-competed for binding to monomeric gp120, indicating the topological proximity of their epitopes, however, the nonneutralizing CD4BS MAbs did not interfere with the neutralization activity of MAb b12 the nonneutralizing MAbs partially competed with b12 for Env binding of the surface of Env-transfected cells this suggests Env has two categories of binding site for CD4BS MAbs, one recognized by both b12 and nonneutralizing CD4BS MAbs, the other is recognized by only b12 Ab-gp120 interactions based on the use of monomeric gp120 or Env-transfected cells do not predict the outcome of HIV-1 neutralization assays, and they should be interpreted with caution. Herrera et al. [2003]
- HT5: Called 205-43-1: To determine the antigenicity of virus killed by thermal and chemical inactivation, retention of conformation-dependent neutralization epitopes was examined, and exposure of CD4BS epitopes was found to be enhanced (MAbs IgG1b12, 205-46-9, and 205-43-1) binding to 2G12 and 447-52D epitopes was essentially unaltered the 17b CD4i epitope was also exposed. Grovit-Ferbas et al. [2000]
- HT5: HT5, HT6, and HT7 are also known as 205-43-1, 205-42-15, and 205-46-9, respectively. Rank order of CD4BS antibodies oligomer binding is IgG1b12 = 2G6 = 205-46-9 > 205-43-1 = 205-42-15 > 15e = 21h = F91, and the only thing notably distinguishing about neutralizing IgG1b12 is that it depends on residues in V2. Fouts *et al.* [1998]
- HT5: HT5 and HT6 bind JRSF oligomer but with low affinity, and are not neutralizing conclusions of this paper contrast with Parren *et al.* [1998a] Fouts *et al.* [1998]. Fouts *et al.* [1998]; Parren *et al.* [1998a]
- HT5: MAbs IgG1b12, HT5, HT6, and HT7 cross-compete for binding to monomeric gp120, bind equally well, inhibit gp120-sCD4 interactions, but only IgG1b12 neutralizes JRFL. Fouts *et al.* [1997]
- HT5: Despite highly cross-reactive binding to many primary and T-cell adapted viral strains, only weakly neutralizes IIIB and MN. Moore et al. [1995a]
- HT5: 205-46-9 was cross-reactive across clades A-F, 205-43-1 very cross-reactive but not quite as extensive 205-46-9. Moore *et al.* [1994b]

**No.** 1080

**MAb ID** HT6 (205-42-15)

**HXB2 Location** Env **Author Location** gp120

Epitope

Neutralizing L (weak)

Immunogen HIV-1 infection

Species (Isotype) human

**Ab Type** gp120 CD4BS

tial glycosylation site. The affinity for sCD4 was unchanged Research Contact Ciba-Geigy AG Basel, Switzerland, and

Tanox Biosystems, Houston, Texas

**References** Pugach *et al.* 2004; Gorny & Zolla-Pazner 2004; Herrera *et al.* 2003; Parren *et al.* 1998a;

Fouts *et al.* 1998; Fouts *et al.* 1997; Moore

et al. 1995a; Moore et al. 1994b

**Keywords** antibody binding site definition and exposure, antibody interactions, reversion, viral fitness, review, subtype comparisons, variant cross-recognition or cross-neutralization

- HT6: Called 205-42-15: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- HT6: Called 205-42-15: A primary isolate, CC1/85, was passaged 19 times in PBMC and gradually acquired increased sensitivity to FAb b12 and sCD4 that was attributed to changes in the the V1V2 loop region, in particular the loss of a potential glycosylation site. The affinity for sCD4 was unchanged in the monomer, suggesting that the structural impact of the change was manifested at the level of the trimer. The passaged virus, CCcon19, retained an R5 phenotype and its neutralization susceptibility to other Abs was essentially the same as CC1/85. Three CD4BS MAbs, 205-46-9, 205-42-15, and 205-43-1, did not neutralize either the primary or passaged variant. Pugach *et al.* [2004] (reversion, viral fitness, variant cross-recognition or cross-neutralization)
- HT6: Called 205-42-15: CD4BS MAbs b12 (neutralizing) and 205-42-15, 204-43-1, 205-46-9 (nonneutralizing) all cross-competed for binding to monomeric gp120, indicating the topological proximity of their epitopes, however, the nonneutralizing CD4BS MAbs did not interfere with the neutralization activity of MAb b12 the nonneutralizing MAbs partially competed with b12 for Env binding of the surface of Envtransfected cells this suggests Env has two categories of binding site for CD4BS MAbs, one recognized by both b12 and nonneutralizing CD4BS MAbs, the other is recognized by only b12 Ab-gp120 interactions based on the use of monomeric gp120 or Env-transfected cells do not predict the outcome of HIV-1 neutralization assays, and they should be interpreted with caution. Herrera *et al.* [2003] (antibody binding site definition and exposure, antibody interactions)
- HT6: HT5, HT6, and HT7 are also known as 205-43-1, 205-42-15, and 205-46-9, respectively. Fouts et al. [1998]
- HT6: HT5 and HT6 bind JRSF oligomer but with low affinity, and are not neutralizing conclusions of this paper contrast with Parren *et al.* [1998a] Fouts *et al.* [1998]. Fouts *et al.* [1998]; Parren *et al.* [1998a] (antibody binding site definition and exposure)

- HT6: MAbs IgG1b12, HT5, HT6, and HT7 cross-compete for binding to monomeric gp120, bind equally well, inhibit gp120sCD4 interactions, but only IgG1b12 neutralizes JRFL. Fouts et al. [1997] (antibody binding site definition and exposure, antibody interactions)
- HT6: Despite highly cross-reactive binding to many primary and T-cell adapted viral strains, only weakly neutralizes IIIB and MN. Moore et al. [1995a] (variant cross-recognition or cross-neutralization)
- HT6: 205-46-9 was cross-reactive across clades A-F, 205-43-1 was not quite as extensively cross-reactive. Moore et al. [1994b] (subtype comparisons)

No. 1081

**MAb ID** HT7 (205-46-9)

**HXB2 Location** Env Author Location gp120

**Epitope** 

Neutralizing L (IIIB)

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

Research Contact Ciba-Geigy AG (Basel, Switzerland), and Tanox Biosystems, Houston, Texas

> References Pugach et al. 2004; Gorny & Zolla-Pazner 2004; Herrera et al. 2003; Grovit-Ferbas et al. 2000; Parren et al. 1998a; Fouts et al. 1998; Fouts et al. 1997; Moore et al. 1995a; Moore et al. 1994b

Keywords antibody binding site definition and exposure, assay standardization/improvement, reversion, viral fitness, review, subtype comparisons, variant cross-recognition or crossneutralization

- HT7: Also called 205-46-9. This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- HT7: Called 205-46-9: A primary isolate, CC1/85, was passaged 19 times in PBMC and gradually acquired increased sensitivity to FAb b12 and sCD4 that was attributed to changes in the V1V2 loop region, in particular the loss of a potential glycosylation site. The affinity for sCD4 was unchanged in the monomer, suggesting that the structural impact of the change was manifested at the level of the trimer. The passaged virus, CCcon19, retained an R5 phenotype and its neutralization susceptibility to other Abs was essentially the same as CC1/85. Three CD4BS MAbs, 205-46-9, 205-42-15, and 205-43-1, did not neutralize either the primary or passaged variant. Pugach et al. [2004] (reversion, viral fitness, variant cross-recognition or cross-neutralization)
- HT7: Called 205-46-9 CD4BS MAbs b12 (neutralizing) and 205-42-15, 204-43-1, 205-46-9 (nonneutralizing) all crosscompeted for binding to monomeric gp120, indicating the topological proximity of their epitopes, however, the nonneutralizing CD4BS MAbs did not interfere with the neutralization activity of MAb b12 – the nonneutralizing MAbs partially competed with b12 for Env binding of the surface of Env- Research Contact Jackie Cordell and C. Dean transfected cells - this suggests Env has two categories of binding site for CD4BS MAbs, one recognized by both b12 and

nonneutralizing CD4BS MAbs, the other is recognized by only b12 – Ab-gp120 interactions based on the use of monomeric gp120 or Env-transfected cells do not predict the outcome of HIV-1 neutralization assays, and they should be interpreted with caution. Herrera et al. [2003] (antibody binding site definition and exposure)

- HT7: Called 205-46-9. To determine the antigenicity of virus killed by thermal and chemical inactivation, retention of conformation-dependent neutralization epitopes was examined, and exposure of CD4BS epitopes was found to be enhanced (MAbs IgG1b12, 205-46-9, and 205-43-1) - binding to 2G12 and 447-52D epitopes was essentially unaltered – the 17b CD4i epitope was also exposed. Grovit-Ferbas et al. [2000] (antibody binding site definition and exposure)
- HT7: Called 205-46-9. HT5, HT6, and HT7 are also known as 205-43-1, 205-42-15, and 205-46-9, respectively. Binds JRSF oligomer with high affinity as does IgG1b12, but IgG1b12 is neutralizing, 205-46-9 is not - conclusions of this paper contrast with Parren98 - authors propose a model where 205-46-9 and 2G6 may inhibit CD4 binding, but cause a conformational shift which enhances CCR5 binding and thus counteracts the neutralizing effect - rank order of CD4BS antibodies oligomer binding is IgG1b12 = 2G6 = 205-46-9 > 205-43-1 = 205-42-15> 15e = 21h = F91, and the only thing notably distinguishing about neutralizing IgG1b12 is that it depends on residues in V2. Fouts et al. [1998] (assay standardization/improvement)
- HT7: Binds JRSF oligomer with high affinity, at least as high as IgG1b12, but IgG1b12 is neutralizing, H7 is not – conclusions of this paper contrast with Parren et al. [1998a] - authors propose a model where H7 may inhibit CD4 binding, but cause a conformational shift which enhances CCR5 binding and thus counteracts the neutralizing effect Fouts et al. [1998]. Fouts et al. [1998]; Parren et al. [1998a] (antibody binding site definition and exposure)
- HT7: MAbs IgG1b12, HT5, HT6, and HT7 cross-compete for binding to monomeric gp120, bind equally well, inhibit gp120sCD4 interactions, but only IgG1b12 neutralizes JRFL. Fouts et al. [1997] (antibody binding site definition and exposure)
- HT7: Despite highly cross-reactive binding to many primary and T-cell adapted viral strains, only neutralizes IIIB well, with sporadic weak neutralization of other isolates. Moore et al. [1995a] (variant cross-recognition or cross-neutralization)
- HT7: 205-46-9 was cross-reactive across clades A-F, 205-43-1 was cross-reactive, but not quite as extensive. Moore et al. [1994b] (subtype comparisons)

**No.** 1082

**MAb ID** ICR 39.13g (ICR39.13g, 39.13g)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

**Species (Isotype)** rat (IgG2b)

Ab Type gp120 CD4BS

References Vella et al. 2002; Peet et al. 1998; Klasse & Sattentau 1996; Armstrong & Dimmock 1996; McKeating et al. 1996; Beretta & Dalgleish 1994; McLain & Dimmock 1994; Klasse et al. 1993a; Thali et al. 1993; Moore & Ho 1993; McKeating et al. 1993b; McKeating et al. 1992c; McKeating et al. 1992a; Cordell et al. Research Contact J. Cordell and C. Dean

- ICR 39.13g: UK Medical Research Council AIDS reagent: ARP390.
- ICR 39.13g: Called ARP390/391, but no such entry was found at the UK Medical Research Council AIDS reagent web site: Herpesvirus saimiri-immortalized CD4+ T lymphocytes (HVS T cells) were used to isolate virus and perform HIV-1 neutralization assays, and compared with a standard PBMC protocol neutralization sensitivities to a panel of MAbs and to homologous or heterologous plasma/sera were similar for HVS T cells (CN-2 cells) and PBMCs. Vella et al. [2002]
- ICR 39.13g: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic – these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind - ICR 39.13g was not affected by V3 serine substitutions - mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]
- ICR 39.13g: Post-attachment neutralization mechanism, in contrast to MAb 39.3b. Armstrong & Dimmock [1996]
- ICR 39.13g: Variants of LAI have differing neutralization susceptibility to 39.13g. Klasse & Sattentau [1996]
- ICR 39.13g: Called 39.13g Neutralizes HXB2, but fails to neutralize chimeric virus with gp120 from primary isolates in an HXB2 background. McKeating et al. [1996]
- ICR 39.13g: Kinetics of neutralization studied no lag for 39.3b, while ICR 39.13g and ICR 41.1i have lags of 5 and 15 min respectively - mediates neutralization with 2.3 molecules of IgG. McLain & Dimmock [1994]
- ICR 39.13g: The gp41 mutation 582(Ala to Thr) results in conformational changes in gp120 that confer neutralization resistance to a class of conformation sensitive neutralizing MAbs – ICR 39.13g required moderately higher concentrations to neutralize the mutant than wild type. Klasse et al. [1993a]
- ICR 39.13g: Neutralization activity against HXB10, RF, SF-2 and MN strains of HIV-1. McKeating et al. [1993b]
- ICR 39.13g: Conformational, does not bind denatured gp120 weak neutralization of IIIB - strong inhibition of HIV+ human sera binding to IIIB gp120. Moore & Ho [1993]
- ICR 39.13g: Strongly inhibits CD4 inducible MAb 48d. Thali et al. [1993]
- ICR 39.13g: Binds to a conformational epitope involved in CD4 binding - exerts a synergistic effect in combination with V3 directed MAbs. McKeating et al. [1992a]
- ICR 39.13g: Cross-competes with MAbs ICR 39.3b and 15e. Cordell et al. [1991]

No. 1083

**MAb ID** ICR 39.3b (39.3, 39.3b, ICR39.3b)

**HXB2 Location** Env **Author Location** gp120 **Epitope** 

Neutralizing L Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2b)

Ab Type gp120 CD4BS

References Wyatt et al. 1998; Jeffs et al. 1996; Armstrong & Dimmock 1996; McLain & Dimmock 1994; Moore et al. 1993b; Moore & Ho 1993; McKeating et al. 1992c; Cordell et al. 1991

- ICR 39.3b: also known as 39.3, 39.3b and ICR39.3b.
- ICR 39.3b: UK Medical Research Council AIDS reagent: ARP391.
- ICR 39.3b: Called 39.3 summary of the implications of the crystal structure of the core of gp120 bound to CD4 and 17b with what is known about mutations that reduce NAb binding probable mechanism of neutralization by CD4BS Ab is direct interference with CD4 binding. Wyatt et al. [1998]
- ICR 39.3b: Neutralizes only if the antibody is added prior to the attachment of the virus to the cell, in contrast to 39.13g. Armstrong & Dimmock [1996]
- ICR 39.3b: Called 39.3b increased binding when V1/V2 or V1/V2 and V3 were deleted from gp120. Jeffs et al. [1996]
- ICR 39.3b: Kinetics of neutralization studied no lag for 39.3b, while ICR 39.13g and ICR 41.1i have lags of 5 and 15 min respectively. McLain & Dimmock [1994]
- ICR 39.3b: Conformational, does not bind to denatured IIIB. Moore & Ho [1993]
- ICR 39.3b: Cross-competes with MAbs ICR 39.13g and 15e. Cordell *et al.* [1991]

No. 1084

MAb ID Ia3

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS

References Zwick et al. 2003

**Keywords** antibody interactions

• Ia3: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This is a novel CD4BS Fab first used in this study. Zwick et al. [2003] (antibody interactions)

**No.** 1085 MAb ID Ia7 HXB2 Location Env

**Author Location** gp120

Epitope Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4BS References Zwick et al. 2003 Keywords antibody interactions

• Ia7: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This is a novel CD4BS Fab first used in this study. Zwick et al. [2003] (antibody interactions)

**No.** 1086

MAb ID IgG1b12 (Fab b12, Fab 3B3, MAb IgG1b12,

IgG1-b12, IgG1 b12, IgGB12, b4/12)

**HXB2 Location** Env Author Location gp120

> **Epitope** Subtype B Neutralizing LP

Immunogen HIV-1 infection **Species** (**Isotype**) human ( $IgG1\kappa$ ) Ab Type gp120 CD4BS

Research Contact D. Burton, Scripps Research Institute, La Jolla, CA, also J. Geltowsky and J. Pyati, R. W. Johnson Pharmaceutical Resear

References Zipeto et al. 2005; Wilkinson et al. 2005; Selvarajah et al. 2005; Ren et al. 2005; Raviv et al. 2005; Pinter et al. 2005; Martín-García et al. 2005; Lusso et al. 2005; Luo et al. 2006; Krachmarov et al. 2005; Heap et al. 2005a; Haynes et al. 2005; Gorny et al. 2005; Chen et al. 2005; Safrit et al. 2004; Pugach et al. 2004; Pinter et al. 2004; Pantophlet et al. 2004; Nabatov et al. 2004; McCaffrey et al. 2004; Jeffs et al. 2004; Ferrantelli et al. 2004a; Dacheux et al. 2004; Biorn et al. 2004; Binley et al. 2004; Zwick et al. 2004; Zwick et al. 2003; Pantophlet et al. 2003b; Zhu et al. 2003; Veazey et al. 2003; Montefiori et al. 2003; Kitabwalla et al. 2003; Zhang et al. 2003; Wang 2003; Mascola 2003; Raja et al. 2003; Hart et al. 2003; Ferrantelli et al. 2003; Dev et al. 2003; Cavacini et al. 2003; Binley et al. 2003; Herrera et al. 2003; Pantophlet et al. 2003a; Poignard et al. 2003; Ling et al. 2002; Lewis et al. 2002; Kwong et al. 2002; Gorry et al. 2002; Cavacini et al. 2002; Bures et al. 2002; Liu et al. 2002; Ferrantelli & Ruprecht 2002; Klasse & Sattentau 2002; Zhang et al. 2002; Grundner et al. 2002; Edwards et al. 2002; Xiang et al. 2002b; Vella et al. 2002; Chakrabarti et al. 2002; Xu et al. 2002; Scanlan et al. 2002; Saphire et al. 2002; Yang et al. 2002; Schulke et al. 2002; Sanders et al. 2002; Golding et al. 2002b; Srivastava et al. 2002; Hezareh et al. 2001; Xu et al. 2001; Hofmann-Lehmann et al. 2001; Verrier et al. 2001; Spenlehauer et al. 2001; Zeder-Lutz et al. 2001; Poignard et al. 2001; Parren et al. 2001; Zwick et al. 2001c; Zwick et al. 2001b; Zwick et al. 2001a; York et al. 2001; Yang et al. 2001; Saphire et al. 2001b; Saphire et al. 2001a; Kolchinsky et al. 2001; Si et al. 2001; Park et al. 2000; Nyambi et al. 2000; Ly & Stamatatos 2000; Grovit-Ferbas et al. 2000; Binley et al. 1999; Beddows et al. 1999; Giraud et al. 1999; Montefiori & Evans 1999; Hioe et al. 1999; Jackson et al. 1999; Crawford et al. 1999; Poignard et al. 1999; Stamatatos & Cheng-Mayer 1998; Kropelin et al. 1998; Frankel et al. 1998; Sullivan et al. 1998a; Schonning et al. 1998; Brand et al. 1998; Parren et al. 1998b; Takefman et al. 1998; Fouts et al. 1998; Binley et al. 1998; Connor et al. 1998; Parren et al. 1998a; Mondor et al. 1998; Wyatt et al. 1998; Valenzuela et al. 1998; Parren & Burton 1997; Parren et al. 1997a; Parren et al. 1997b; Boots et al. 1997; Burton & Montefiori 1997; Wyatt et al. 1997; Ugolini et al. 1997; Ditzel et al. 1997; Stamatatos et al. 1997; Moore & Trkola 1997; Kessler II et al. 1997; Li et al. 1997; Fouts et al. 1997; Mo et al. 1997; Schutten et al. 1997; D'Souza et al. 1997; McKeating 1996; Sattentau 1996; Trkola et al. 1996a; Poignard et al. 1996a; Poignard et al. 1996b; Gauduin et al. 1996; Moore & Sodroski 1996; Yang et al. 1997c; Sullivan et al. 1995; Ditzel et al. 1995; Trkola et al. 1995; Parren et al. 1995; Moore & Ho 1995; Moore et al. 1995a; Sattentau 1995; Sattentau et al. 1995; Kessler et al. 1995; Moore et al. 1994b; Burton et al. 1994; Roben et al. 1994; Barbas III et al. 1992; Burton et al. 1991

Keywords antibody binding site definition and exposure, antibody generation, antibody interactions, antibody sequence, variable domain, co-receptor, immunoprophylaxis, neutralization potency, review, structure, subtype comparisons, vaccine antigen design, vaccinespecific epitope characteristics, variant crossrecognition or cross-neutralization

- IgG1b12: Fab b12 was derived from IgG1b12, Fab 3B3 was derived from Fab b12 by random mutagenesis and selected for increased affinity to sgp120 - database note. (antibody generation)
- IgG1b12: UK Medical Research Council AIDS reagent: ARP3065.

- IgG1b12: NIH AIDS Research and Reference Reagent Program: 2640.
- IgG1b12: gp41 and p15E of the porcine endogenous retrovirus (PERV) share structural and functional similarities, and epitopes in the membrane proximal region of p15E are able to elicit NAbs upon immunization with soluble p15E. Rabbits immunized with a VSV recombinant expressing an HIV-1 membrane-proximal external region (MPER) fused to PERV p15E, with a fusion p15E-HIV MPER protein boost, elicited HIV specific NAbs. The IgG1b12 MAb was used as a positive control for neutralization in this study. Luo *et al.* [2006] (vaccine antigen design)
- IgG1b12: The lack of glycosylation sites at residues Asn 295 and Thy 394 within C-clade gp120s generally causes the loss of 2G12 recognition. Introduction of glycans in the subtype C strain HIV-1CN54 at these positions restored 2G12 binding, and addition of just a single glycan partially restored binding (V295N + A394T » V295N > A395T). 2G12 epitope recovery decreased b12 binding. Chen *et al.* [2005]
- IgG1b12: 2909 is a human anti-Env NAb that was selected by neutralization assay and binds to the quaternary structure on the intact virion. ELISA-based competition assays and subsequent mutational analysis determined that the CD4BS and V2 and V3 loops contribute to the 2909 epitope: 2909 binding was inhibited by MAbs 447-52d (anti-V3), 830A (anti-V2), and IgG1b12 (anti-CD4BS) and sCD4. 2909 was not inhibited by MAbs 670, 1418, nor 2G12. Gorny *et al.* [2005]
- IgG1b12: IgG1b12, like the other anti-Env broadly neutralizing MAbs 2F5 and 4E10, binds to auto-antigens and has characteristics of polyspecific autoreactive antibodies. Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. IgG1B12 reacted with ribonucleoprotein, dsDNA, centromere B, and histones, as well as nucleolar and cytoplasmic reactivity in HEp-2 cells. Haynes *et al.* [2005]
- IgG1b12: b12 and the gp41 C-terminal binding MAb SAR1 inhibit HIV-1 infected cell fusion with target cells at comparable levels. Heap *et al.* [2005a]
- IgG1b12: Sera from subtype A infected individuals from Cameroon have antibodies that react strongly with subtype A and subtype B V3 loops in fusion proteins, and neutralize SF162 pseudotypes, while sera from 47 subtype B infected individuals reacted only with subtype B V3s. Sera from Cameroon did not neutralize primary A or B isolates, due to indirect masking by the V1/V2 domain rather than due to loss of the target epitope. Neutralization by Cameroonian sera MAbs was blocked by Clade A and B V3 loop fusion proteins, while NAbs to non-V3 epitopes, 2F5, 2G12, and b12, were not blocked. Krachmarov *et al.* [2005]
- IgG1b12: Called IgG1 b12. The HIV-1 Bori-15 variant was adapted from the Bori isolate for replication microglial cells. Bori-15 had increased replication in microglial cells and a robust syncytium-forming phenotype, ability to use low levels of CD4 for infection, and increased sensitivity to neutralization by sCD4 and 17b. Four amino acid changes in gp120 V1-V2 were responsible for this change. Protein functionality and

- integrity of soluble, monomeric gp120-molecules derived from parental HIV-1 Bori and microglia-adapted HIV-1 Bori-15 was assessed in ELISA binding assays using CD4BS MAbs F105 and IgG1b12, glycan-specific 2G12, and V3-specific 447-52D, and were unchanged. Association rates of sCD4 and 17b were not changed, but dissociation rates were 3-fold slower for sCD4 and 14-fold slower for 17b. Martín-García *et al.* [2005]
- IgG1b12: This study is about the V2 MAb C108g, that is type-specific and neutralizes BaL and HXB2. JR-FL is a neutralization resistant strain; modification of JRFL at V2 positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The modification in V2 also increased neutralization sensitivity to V3 MABs 4117c, 2219, 2191, and 447-52D, but only had minor effects on neutralization by CD4BS MAb 5145A, and broadly neutralizing MAbs IgG1b12, 2G12, and 2F5. gp120 binding to CD4 was inhibited by b12, but not by C108g. Pinter *et al.* [2005]
- IgG1b12: Retrovirus inactivation for vaccine antigen delivery was explored through lipid modification by hydrophobic photoinduced alkylating probe 1.5 iodonaphthylazide (INA). The viral proteins were shown to be structurally intact in the treated non-infectious virus, through the preservation of antibody binding sites for polyclonal anti-gp120 serum, and for broadly neutralizing MAbs 2G12, b12 and 4E10, although the modifications of the lipid disabled viral infection. Raviv *et al.* [2005] (vaccine antigen design)
- IgG1b12: The antibody M2 is specific for a peptide flag inserted into the V4 loop of YU-2, a neutralization resistant variant with a short V4 loop. IgG1b12 and 2F5 could neutralize both the WT YU-2 and the modified variant. The high diversity of V4 suggests it does not play a direct role in receptor binding or viral entry, yet M2, specific for the peptide insert tag, can neutralize the modified virus, demonstrating that neutralizing activity doesn't have to block functionality of the virus. Ren et al. [2005] (neutralization potency)
- IgG1b12: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies in vaccinated rabbits, but GDMR elicited anti-V3 NAbs. Both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. CD4BS MAbs except Fab b12 (b6, b3, F105) did not bind to either GDMR or mCHO. CD4i MAbs (48d, 17b) did not bind even with sCD4. 2G12 had diminished binding to both. V3 MAbs (447-52D, 19b, F245-B4e8 and 39F) bound to the GDMR antigen, but either did not bind or had diminished binding to mCHO. V2 MAb 697-D did not bind to mCHO and had diminished binding to GDMR, while V2 MAb 8.22.2 bound to GDMR but not mCHO. V1/V2/V3 MAb 4KG2, C1-C4 MAb A32, C1-C5 MAb C11, and HIVIG all either did not bind or had significantly diminished binding to both antigen constructs. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)
- IgG1b12: The crystal structure of the Fab fragment from F105 was solved. It has an extended CDR H3 loop, with a Phe at the apex that may recognize the binding pocket of gp120

used by the Phe-42 residue of CD4. The potent NAB IgG1b12 recognizes an overlapping binding site, the main difference is that F105 extends across the interface of the inner and outer domains of gp120 while b12 does not. IgG1b12 also has undergone extensive affinity maturation (45 mutations) while F105 has not (13 mutations) – an average for gp120 MAbs is 22 mutations. Wilkinson *et al.* [2005] (antibody sequence, variable domain, structure)

- IgG1b12: HIV-1 fusion complexes were prepared from cell lines expressing R5 HIV-1 gp120/gp41 and CD4-CCR5. Neutralizing Abs were raised against both R5 (strain BaL) and X4 (strain 213) viruses. IgG1b12 was used to detect gp120/gp41. Zipeto *et al.* [2005] (vaccine antigen design)
- IgG1b12: 93 viruses from different clades were tested for their neutralization cross-reactivity using a panel of HIV antibodies. IgG1b12 neutralized a fraction of viruses from almost every clade, and was more potent that 2F5 and 4E10, particularly against a subset of B clade viruses. Binley et al. [2004] (variant cross-recognition or cross-neutralization, subtype comparisons)
- IgG1b12: Called b12. The peptide 12p1 (RINNIPWSEAMM) inhibits direct binding of YU2 gp120 or Env trimer to CD4, CCR5 and MAb 17b in a concentration-dependent allosteric manner. 12p1 is thought to bind to unbound gp120 near the CD4 binding site, with a 1:1 stoichiometry. 12p1 also inhibited MAb F105 binding. presumably because F105 favors an unactivated conformation, but not 2G12 or b12. The 1:1 stoichiomtry, the fact that the peptide binding site is accessible on the trimer, the non-CD4 like aspect of the binding, and an ability to inhibit viral infection in cell cultures make it a promising lead for therapeutic design. Biorn *et al.* [2004] (antibody binding site definition and exposure)
- IgG1b12: Env sequences were derived from 4 men at primary infection and 4 years later; the antigenicity in terms of the ability to bind to 2G12, 2F5 and IgG1b12 was determined. 2G12 bound primarily to late clones in 3 of the 4 patients, and to both early and late in the other patient. Neither 2F5 nor IgG1b12 showed a difference in binding affinity to early or late envelopes. Dacheux *et al.* [2004]
- IgG1b12: Nabs against HIV-1 M group isolates were tested for their ability to neutralize 6 randomly selected HIV-1 O group strains. IgG1b12 could neutralize some O group strains when used on its own, and quadruple combination of b12, 2F5, 2G12, and 4E10, could neutralize the six Group O viruses tested between 62-97%. Ferrantelli *et al.* [2004a]
- IgG1b12: Called b12. A set of oligomeric envelope proteins were made from six primary isolates for potenial use as vaccine antigens: 92/UG/037 (clade A), HAN2/2 (clade B), 92/BR25/025 (clade C), 92/UG/021 (clade D), 93/BR/029 (clade F) and MVP5180 (clade O). This was one of a panel of MAbs used to explore folding and exposure of well characterized epitopes. The clade C isolate BR25 is apparently misfolded, as conformation-dependent antibodies did not bind to it. b12 bound to clade A, B, D and F HIV-1 primary isolates. Polyclonal sera raised in rabbits against these antigens cross-bound the other antigens, but none of the sera had neutralizing activity. Jeffs et al. [2004] (variant cross-recognition or cross-neutralization)

- IgG1b12: Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The loss of the any of the five glycans, within the V3 loop (GM299 V3), C2 (GM292 C2), C3 (GM329 C3), C4 (GM438 C4), or V5 (GM454 V5) made SF162 become more sensitive to IgG1b12 neutralization. V3 glycans tended to shield V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans shielded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004] (antibody binding site definition and exposure)
- IgG1b12: Fab b12. A set of HIV-1 chimeras that altered V3 net charge and glycosylation patterns in V1V2 and V3, involving inserting V1V2 loops from a late stage primary isolate taken after the R5 to X4 switch, were studied with regard to phenotype, co-receptor usage, and MAb neutralization. The loops were cloned into a HXB2 envelope with a LAI viral backbone. It was observed that the addition of the late-stage isolate V1V2 region and the loss of V3-linked glycosylation site in the context of high positive charge gave an X4 phenotype. R5X4 viruses were more sCD4 and 2G12 neutralization resistant than either R5 or X4, but the opposite pattern was observed for b12. Addition of the late stage V1V2 altered neutralization for both MAbs, but this alteration was reversed with the loss of the V3 glycan. Nabatov *et al.* [2004] (**co-receptor**)
- IgG1b12: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it binds to the neutralizing MAb 2G12. It masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120. Pantophlet *et al.* [2004]
- IgG1b12: Called IgG-b12. V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12 which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Three anti-CD4BS MAbs were tested, including IgG1b12 which neutralizes both JRFL and SF162. The affinities for IgG1b12 and 5145A were similar for both JRFL and SF612, but 1125A bound with 2.5 fold higher affinity to SF162. 5145A and 1125H both preferentially neutralize SF162, but not JRFL, and the CD4BS is more sensitive to neutralization in the context of the SF162 V1V2 loop. This was also true for neutralization by sCD4. Pinter et al. [2004] (variant crossrecognition or cross-neutralization)
- IgG1b12: Called b12. A primary isolate, CC1/85, was passaged 19 times in PBMC and gradually acquired increased

sensitivity to FAb b12 and sCD4 that was attributed to changes in the V1V2 loop region, in particular the loss of a potential glycosylation site. The affinity for sCD4 was unchanged in the monomer, suggesting that the structural impact of the change was manifested at the level of the trimer. The passaged virus, CCcon19, retained an R5 phenotype and its neutralization susceptibility to other Abs was essentially the same as CC1/85. CCcon19 (IC50 0.3) was significantly more senstivie to neutralization by b12 than was CC1/85 (IC50 6.0). Pugach *et al.* [2004]

- IgG1b12: This review discusses research presented at the Ghent Workshop of prevention of breast milk transmission and immunoprophylaxis for HIV-1 in pediatrics (Seattle, Oct. 2002), and makes the case for developing passive or active immunoprophylaxis in neonates to prevent mother-to-infant transmission. Macaque studies have shown that passive transfer of NAb combinations (for example, IgG1b12, 2G12, 2F5, and 4E10) can confer partial or complete protection to infant macaques from subsequent oral SHIV challenge. Safrit et al. [2004]
- IgG1b12: Called IgG1 b12. This paper is a study of the 2F5 NAb complexed to peptide ELDKWAS; the peptide was found to interact with amino acids near the base of the very long (22 residue) CDR 3H region of the Ab, although a Phe at the apex of the loop was also important. The authors suggest that particularly long CDR H3 regions may be a common feature of HIV-1 neuturalizing antibodies there are 22 residues in 2F5's H3, 18 in IgG1b12's H3, and 22 residues in X5's H3. They express concern that because small animals like mice are unable to elicit Ab responses with such long H3s, they may be poor model systems for HIV vaccine studies. Zwick *et al.* [2004] (antibody sequence, variable domain)
- IgG1b12: The SOS mutant envelope protein introduces a covalent disulfide bond between gp120 surface and gp41 transmembrane proteins into the R5 isolate JR-FL by adding cysteines at residues 501 and 605. Pseudovirions bearing this protein bind to CD4 and co-receptor bearing cells, but do not fuse until treatment with a reducing agent, and are arrested prior to fusion after CD4 and co-receptor engagement. IgG1b12 neutralized SOS and WT proteins comparably, and neither IgG1b12 nor the Fab b12 could neutralize well post-attachment, consistent with the notion that theb12 binding site would be blocked upon cellular binding. Binley et al. [2003]
- IgG1b12: Called 1b12. The MAb B4e8 binds to the base of the V3 loop, neutralizes multiple primary isolates and was studied for interaction with other MAbs. CD4BS MAb IgG1b12 had no effect on B4e8 binding. Cavacini *et al.* [2003]
- IgG1b12: A sCD4-17b single chain chimera was made that can bind to the CD4 binding site, then bind and block co-receptor interaction. This chimeric protein is a very potent neutralizing agent, more potent than IgG1b12, 2G12 or 2F5 against Ba-L infection of CCR5-MAGI cells. It has potential for prophylaxis or therapy. Dey *et al.* [2003]
- IgG1b12: Four newborn macaques were challenged with pathogenic SHIV 89.6 and given post exposure prophylaxis using a combination of NAbs 2F5, 2G12, 4E10 and IgG1b12. 2/4 treated animals did not show signs of infection, and 2/4 macaques maintained normal CD4+ T cell counts and had a lower delayed peak viremia compared to the controls. Ferrantelli *et al.* [2003] (immunoprophylaxis)

- IgG1b12: Called b12 CD4BS MAbs b12 (neutralizing) and 205-42-15, 204-43-1, 205-46-9 (nonneutralizing) all cross-competed for binding to monomeric gp120, indicating the topological proximity of their epitopes, however, the nonneutralizing CD4BS MAbs did not interfere with the neutralization activity of MAb b12 the nonneutralizing MAbs partially competed with b12 for Env binding of the surface of Env-transfected cells this suggests Env has two categories of binding site for CD4BS MAbs, one recognized by both b12 and nonneutralizing CD4BS MAbs, the other is recognized by only b12 Ab-gp120 interactions based on the use of monomeric gp120 or Env-transfected cells do not predict the outcome of HIV-1 neutralization assays, and they should be interpreted with caution. Herrera *et al.* [2003] (antibody binding site definition and exposure)
- IgG1b12: MAbs IgG1b12, 2G12, 2F5 and 4E10 were tested for their ability to neutralize two primary HIV-1 clade A isolates (UG/92/031 and UG/92/037) and two primary HIV-1 clade D isolates (UG/92/001 and UG/92/005). 4E10 demonstrated the most potent cross-neutralization activity. Quadruple administration of MAbs IgG1b12, 2G12, 2F5, and 4E10 induced strong synergistic neutralization of 4 clade A isolates (UG/92/031, UG/92/037, RW/92/020 and RW/92/025) as well as 5 clade D isolates (UG/92/001,UG/9/005, /93/086/RUG/94/108, UG/94/114). The authors note this combination of 4 MAbs neutralizes primary HIV A, B, C, and D isolates. Kitabwalla *et al.* [2003] (variant cross-recognition or cross-neutralization, subtype comparisons)
- IgG1b12: This review dicusses the importance and function of protective antibody responses in animal model studies in the context of effective vaccine development. SHIV models have shown protection using high levels of MAbs can prevent infection, and partial protection that can influence disease course can be obtained from modest levels of NAbs. SHIV challenges studies conducted with infusions of combinations of MAbs b12, 2G12, and 2F5 are reviewed. Mascola [2003] (review)
- IgG1b12: AC10 is a subject who was given treatment early after infection, and had a viral rebound after cessation of therapy, which then declined to a low level. The polyclonal sera from AC10 could potently neutralize the rebound virus, and NAb escape followed with a neutralizing response against the escape variant and subsequent escape from that response. Viral loads remained low in this subject despite escape. The rebound isolate that was potently neutralized by autologous sera was not particularly neutralization sensitive, as it resisted neutralization by sCD4 and MAbs IgG1b12, 2G12 and 2F5, and was only moderately sensitive to sera from other HIV+ individuals that had high titers of NAbs to TCLA strains. Montefiori *et al.* [2003]
- IgG1b12: Called b12 Alanine scanning mutagenesis was used to compare substitutions that affected anti-CD4BS NAb b12 binding to those that affect binding of sCD4 and two non-neutralizing anti-CD4BS Abs b3 and b6 while the epitope maps overlapped, there were some differences observed binding of CD4 was never enhanced, indicating it had evolved to be optimal rec gp120s were engineered to contain combinations of Alanine substitutions that enhanced b12 binding, and while binding of b12 to these gp120 monomers was generally maintained or increased, binding by five non-neutralizing

anti-CD4bs MAbs (b3, b6, F105, 15e, and F91) was reduced or completely abolished – 2G12 binding was largely unperturbed, indicating these proteins were not grossly misfolded – for twelve mutants, b12 neutralization sensitivity and affinity correlated, but for five mutants neutralization efficiency was maintained or increased despite a decrease in affinity suggesting that the substitutions that influence b12 binding to the monomer are different than those that impact neutralization sensitivity to the trimer. Pantophlet *et al.* [2003a] (antibody binding site definition and exposure)

- IgG1b12: This paper describes an attempt to engineer a gp120 molecule that would focus the immune response onto the IgG1b12 epitope. Four Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with seven N-linked glycosylation site sequons and this combination minimized the binding of non-neutralizing MAbs. b12 affinity was lowered, and binding of non-neutralizing MAbs was knocked out. C1 and C5 regions were then removed to eliminate the epitopes for MAbs against these regions, but these also diminished IgG1b12 binding. Pantophlet *et al.* [2003b] (vaccine antigen design)
- IgG1b12: Virion capture assays are not a good predictor of neutralization, and the presentation of epitopes using this assay seems to be different from that of functional Envelope spikes on primary isolates F105 and b6 could efficiently block the b12-mediated capture of infectious virions in a virus capture, but did not inhibit b12 neutralization b12 was potent at neutralizing the three primary virions JR-CSF, ADA, and 89.6, but anti-V3 Abs 447-52D and 19b, which did not neutralize JR-CSF and ADA captured amounts of p24 equal to or higher than the amounts captured by the neutralizing Ab b12. Poignard *et al.* [2003] (neutralization potency)
- IgG1b12: This paper shows that binding of CD4BS MAbs to Env blocks the conformational shift that allows co-receptor CCR5 binding and CD4-independent mediated cell fusion. CD4BS MAbs F105, 15e, and IgG1b12 as well as their Fab fragments inhibited CD4-independent binding of the V1/V2 loop-deleted gp120 glycoproteins of R5 HIV-1 isolates ADA, YU2 and JR-FL and to CCR5 in a concentration dependent manner. CD4BS MAbs IgG1b12, F91 and F105 and their Fab counterparts (except for C11, used as a negative control) inhibited CD4-independent JR-FL and YU-2 gp120-CCR5 binding to CCR5-expressing Cf2Th cells and syncytium formation. Raja et al. [2003] (antibody binding site definition and exposure)
- IgG1b12: Called b12. The NAb b12 was administered locally to the vagina in macaques and could protect against subsequent vaginal infection with SHIV-162P4. This NAb model of a topical microbicide was dose dependence, and was effective for up to 2 hours after administration. Veazey *et al.* [2003] (immunoprophylaxis)
- IgG1b12: Called b12. Review of current neutralizing antibodybased HIV vaccine candidates and strategies of vaccine design. Strategies for targeting of the epitopes for NAbs 2F5, 2G12, 4E10, b12, and Z13 are described. Wang [2003] (review)
- IgG1b12: Called b12. The Fab m18 was selected from a human phage display library by a new method called sequential antigen panning (SAP), using a series of antigens to screen the library to pick broadly cross-reactive isolates. The ability to block cell

- mediated fusion by m17 was compared to Fabs X5 and b12 for a clade A, CRF01 EA, G, and 6 clade B isolates, and the inhibitory activity of m18 was slightly lower but comparable to neutralizing Fabs b12 and X5. Zhang *et al.* [2003]
- IgG1b12: The HIV-1 primary isolate DH012 has preserved the epitopes for the MAbs IgG1b12, 2G12, 17b, however natural DH012 infection in chimpanzees and DH012 gp120 vaccination in guinea pigs does not give rise to Abs against these epitopes. Zhu et al. [2003] (vaccine antigen design)
- IgG1b12: 4KG5, a single-chain Fv (scFv), reacts with a conformational epitope that is formed by the V1, V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Denaturation of gp120 abolished binding of 4KG5 and Fab b12. Additionally, binding of 4KG5 was abrogated when any of the V1, V2 or V3 loops were deleted. Of a panel of Abs tested, only NAb b12 enhanced 4KG5 binding to gp120 JR-FL. MAbs to the following regions diminished or abroated binding: V2 loop MAbs (G3-4, G3-136), V3 loop MAbs (19b, 447-52D, hNM01, AH48, loop2, F425 B4e8, 694-88D), V3-C4 (G3-299, G3-42, G3-519, G3-537), CD4BS (b6, b3, F91, F105, 15e, L33, 1008-D, 654-30D, 559-64D, 1027-30D, Ia3, Ia7, FG39, Fbb14). MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1, V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. 4KG5 did not enhance IgG1b12 neutralization. Zwick et al. [2003] (antibody interactions)
- IgG1b12: IgG1b12 neutralized many South African (5/8) and Malawian (4/8) clade C primary HIV-1 isolates, being more effective than 2F5 which neutralized only two Malawian and no South African isolates. 2G12 did not neutralize any of the 16 isolates. Bures *et al.* [2002] (variant cross-recognition or cross-neutralization, subtype comparisons)
- IgG1b12: This study examined Ab interactions, binding and neutralization with a B clade R5 isolate (92US660) and R5X4 isolate (92HT593). Abs generally bound and neutralized the R5X4 isolate better than the R5 isolate. Anti-gp41 MAb F240 enhanced the binding of CD4BS MAbs IgG1b12 and F105 to both R5X4 and R5 isolates, but had no effect on neutralization. Anti-V3 MAb B4a1 increased CD4BS MAbs IgG1b12 and F105 to R5X4 virions, but only IgG1b12 binding was increased by B4a1 to the R5 isolate, and neutralization was not impacted. Cavacini *et al.* [2002] (antibody interactions)
- IgG1b12: A modified gp140 (gp140ΔCFI), with C-term mutations intended to mimic a fusion intermediate and stabilize trimer formation, retained antigenic conformational determinants as defined by binding to CD4 and to MAbs 2F5, 2G12, F105, and b12, and enhanced humoral immunity without diminishing the CTL response in mice injected with a DNA vaccine. Chakrabarti *et al.* [2002]
- IgG1b12: Truncation of the gp41 cytoplasmic domain of X4, R5, and X4R5 viruses forces a conformation that more closely resembles the CD4 bound state of the external Envelope, enhancing binding of CD4i MAbs 17b and 48d and of CD4BS MAbs F105, b12, and in most cases of glycosylation site dependent MAb 2G12 and the anti-gp41 MAb 246D – in contrast, binding of the anti-V2 MAb 697D and the anti-V3 MAb 694/98D were not affected – viruses bearing the truncation

were more sensitive to neutralization by MAbs 48d, b12, and 2G12 – the anti-C5 MAb 1331A was used to track levels of cell surface expression of the mutated proteins. Edwards *et al.* [2002] (antibody binding site definition and exposure)

- IgG1b12: Review of NAbs that notes IgG1b12 is a recombinant IgG1 from a phage displayed Fab generated against gp120 from a B clade infected individual, that it binds the CD4BS, that alone or in combination with other MAbs it can protect some macaques against SHIV infection, and that it has strong ADCC activity. Ferrantelli & Ruprecht [2002] (review)
- IgG1b12: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion the preincubation 31.5 C step did not alter the inhibitory activity of neutralizing Abs anti-gp41 2F5, or anti-gp120 2G12, IG1b12, 48d, and 17b. Golding *et al.* [2002b]
- IgG1b12: HIV-1 gp160ΔCT (cytoplasmic tail-deleted) proteoliposomes (PLs) containing native, trimeric envelope glycoproteins from R5 strains YU2 and JRFL, and X4 strain HXBc2, were made in a physiologic membrane setting as candidate immunogens for HIV vaccines—2F5 bound to gp160ΔCT with a reconstituted membrane ten-fold better than the same protein on beads—anti-CD4BS MAbs IgG1b12 and F105, A32 (C1-C4), C11 (C1-C5), and 39F (V3) MAbs bound gp160ΔCT PLs indistinguishably from gp160ΔCT expressed on the cell surface—non-neutralizing MAbs C11 and A32 bound with lower affinity than NAb IgG1b12—the MAb 17b was sCD4 inducible on gp160ΔCT PL. Grundner et al. [2002]
- IgG1b12: A broad review of NAbs that mentions IgG1b12 as an example of a NAb that does not alter the conformation of gp120, but interferes with CD4 binding. Klasse & Sattentau [2002] (review)
- IgG1b12: Called b12. Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. Enthalpy and entropy changes were divergent, but compensated. CD4 and MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs (17b, 48d, 1.5e, b6, F105 and F91) had large entropy contributions to free energy of binding to the gp120 monomer (mean: 26.1 kcal/mol, range 18.6-31.5), but the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. The high values suggest surface burial or protein folding and ordering of amino acids upon binding. NAb 2G12 had an entropy value of -1.6. These results suggest that while the trimeric Env complex has four surfaces, a nonneutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding that is not faced by other antigp120 antibodies. Kwong et al. [2002] (structure)
- IgG1b12: Recombinant adeno-associated virus was used to deliver the IgG1b12 gene into mice by injection. IgG1b12 was expressed in these mice for over 6 months after the primary

- injection. This strategy allows for predetermined Ab specificity, and could ultimately be used with synergistic Ab combinations. Lewis *et al.* [2002]
- IgG1b12: A pseudotyping assay showed that an X4 V3 loop peptide could enhance infectivity of X4 virus, R5 and R5X4 V3 loops peptides could enhance infectivity of an R5 virus, and R5X4 peptides could enhance infectivity of an R5X4 virus. Neither R5 nor R5X4 peptides influenced binding of CD4BS MAbs F105 and Ig1Gb12, but did increase binding of CD4i MAb 17b. Ling et al. [2002]
- IgG1b12: Review of NAbs that discusses mechanisms of neutralization, passive transfer of NAbs and protection in animal studies, and vaccine strategies. Liu et al. [2002] (review)
- IgG1b12: Deglycosylation of gp120 does not significantly affect IG1b12 binding, in contrast to MAB 2G12. Sanders et al. [2002]
- IgG1b12: The crystal structure of IgG1b12 is resolved and is the first structure of an intact human Ab with an ordered, full length hinge the structure is extremely asymmetric and flexible with an antigen-binding site that has an unusually long CDR H3 region with a ten residue insertion that projects above the rest of the antigen-binding site this loop may be required for recognition of the recessed CD4 binding site of gp120. Saphire *et al.* [2002] (antibody binding site definition and exposure, antibody sequence, variable domain, structure)
- IgG1b12: Alanine scanning mutagenesis used in conjunction with competition and replacement studies of N-linked carbohydrates and sugars suggest that the 2G12 epitope is formed from mannose residues contributed by the glycans attached to N295 and N332, with the other N-linked carbohydrates in positions N339, N386, and N392 playing a role in maintaining conformation relevant to 2G12 binding N295A and N332A mutants showed essentially unchanged anti-CD4BS NAb b12 binding affinities, while N339A, N386A and N392A mutants displayed significantly lowered b12 affinity, presumably due to conformational changes. Scanlan *et al.* [2002] (antibody binding site definition and exposure)
- IgG1b12: Ab binding characteristics of SOS gp140 were tested using SPR and RIPA SOS gp140 is gp120-gp41 bound by a disulfide bond NAbs 2G12, 2F5, IgG1b12, CD4 inducible 17b, and 19b bound to SOS gp140 better than uncleaved gp140 (gp140unc) and gp120 non-neutralizing MAbs 2.2B (binds to gp41 in gp140unc) and 23A (binds gp120) did not bind SOS gp140 SOS gp140-2F5-IgG1b12 formed multiple ring structures composed of two SOS gp140 proteins bridged by two Ab molecules, while 2F5 and 2G12 formed extended chains rather than closed rings. Schulke *et al.* [2002]
- IgG1b12: Oligomeric gp140 (o-gp140) derived from R5 primary isolate US4 was characterized for use as a vaccine reagent

   antigen capture ELISA was used to compare the antigenicity of gp120 and o-gp140 using a panel of well characterized MAbs Abs directed against the CD4 binding site (IgGCD4 and IgG1b12) reacted slightly more strongly with the gp120 monomer than with the oligomer, as did sCD4. Srivastava et al. [2002]
- IgG1b12: Called ARP3065: Herpesvirus saimiri-immortalized CD4+ T lymphocytes (HVS T cells) were used to isolate virus and perform HIV-1 neutralization assays, and compared with a standard PBMC protocol neutralization sensitivities to a

panel of MAbs and to homologous or heterologous plasma/sera were similar for HVS T cells (CN-2 cells) and PBMCs. Vella *et al.* [2002] (**neutralization potency**)

- IgG1b12: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations 375 S/W seems to favor a conformation of gp120 closer to the CD4-bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang et al. [2002b] (antibody binding site definition and exposure, neutralization potency)
- IgG1b12: Passive immunization of neonate macaques with a combination of F105+2G12+2F5 conferred complete protection against oral challenge with SHIV-vpu+ or the combination b12+2G12+2F5 conferred partial protection against SHIV89.6 such combinations may be useful for prophylaxis at birth and against milk born transmission the synergistic combination of IgG1b12, 2G12, 2F5, and 4E10 neutralized a collection of HIV clade C primary isolates. Xu *et al.* [2002]
- IgG1b12: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin—stabilized oligomer gp140Δ683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and C11, A32, and 30D which did not bind the stabilized oligomer. Yang *et al.* [2002]
- IgG1b12: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang *et al.* [2002] (antibody binding site definition and exposure)
- IgG1b12: Called IgG1 b12. IgG1b12 induces strong ADCC and CDC cytoxicity of HIV-1 infected cells. A panel of mutants in the Fc region of IgG1b12 was generated. K322A reduced ADCC binding of Fc $\gamma$ R and abolished complement-dependent cytotoxicity (CDC) and C1q binding. L234A plus L235 in the lower hinge region of the IgG1 heavy chain abolished both Fc $\gamma$ R and C1q binding and ADCC and CDC. These mutants did not impact IgG1b12's ability to neutralize virus. Hezareh et al. [2001]
- IgG1b12: A combination of MAbs IgG1b12, 2F5, and 2G12 was given postnatally to four neonates macaques that were then challenged with highly pathogenic SHIV89.6P one of

the four infants remained uninfected after oral challenge, two infants had no or a delayed CD4(+) T-cell decline – the most potent combination included IgG1b12, which alone does not alone neutralize SHIV89.6P. Hofmann-Lehmann *et al.* [2001]

- IgG1b12: Mutations in two gloosylation sites in the V2 region of HIV-1 ADA at positions 190 and 197 (187 DNTSYRLINC-NTS) cause the virus to become CD4-independent and able to enter cells through CCR5 alone these same mutations tended to increase the neutralization sensitivity of the virus, except the mutation 197 S/R which resulted in a carbohydrate addition to 195 N that disrupts the IgG1b12 binding site. Kolchinsky *et al.* [2001]
- IgG1b12: Intravenous passive transfer of MAb b12 provides dose-dependent protection from infection to macaques vaginally challenged with the R5 virus SHIV(162P4) the primary isolate HIV-1SF162 is neutralized 90% (IC90) by b12 at 2 μg/ml, and SHIV162P4, derived from HIV-1SF162, was neutralized by 90% at 2 μg/ml in PHA-activated PBMC from rhesus macaques the 90% neutralization titers achieved in three groups of animals that were given 25-, 5-, and 1-mg/kg doses were approximately 1:400, 1:80, and 1:16, respectively the half-life of IgG1 b12 in plasma was about 1 week, but while the peak b12 plasma concentration was immediately after the infusion, the peak vaginal fluid concentration was 7-14 days later. Parren *et al.* [2001]
- IgG1b12: Structural aspects of the interaction of neutralizing Abs with HIV-1 Env are reviewed - Env essentially has three faces, one is largely inaccessible on the native trimer, and two that exposed but have low immunogenicity on primary viruses - neutralization is suggested to occur by inhibition of the interaction between gp120 and the target cell membrane receptors as a result of steric hindrance and it is noted that the attachment of approximately 70 IgG molecules per virion is required for neutralization, which is equivalent to about one IgG molecule per spike – the 2G12, 17b and b12 epitopes are discussed in detail - the structure of CD4-bound gp120 reveals features that HIV has evolved to escape anti-CD4BS Abs like IgG1b12 despite profound functional constraints – CD4BS Abs must first access the CD4 binding site, deeply recessed within the gp120 core, and the Fab of an Ab molecule is "wider" than CD4, and in addition the binding site is flanked by variable and glycosylated regions. Poignard et al. [2001]
- IgG1b12: This paper describes the technical aspects of the crystallization of b12 at a resolution of 2.7 angstroms with all 12 Ig domains resolved. Saphire *et al.* [2001a]
- IgG1b12: This paper describes the biological implications of the crystal structure of b12 a remarkable feature of this antibody is a long protruding finger-like CDR H3 that can dock in the recessed CD4-binding site a contact residues in gp120 are modeled, with numbering based on the variable loop-deleted crystal structure of gp120. Saphire *et al.* [2001b]
- IgG1b12: SHIV-HXBc2 is a neutralization sensitive non-pathogenic virus, and several *in vivo* passages through monkey's yielded highly pathogenic SHIV KU-1 HXBc2 and the KU-1 clone HXBc2P3.2 differ in 12 amino acids in gp160 substitutions in both gp120 and gp41 reduced the ability of sCD4, IgG1b12, F105 and AG1121 to Env achieve saturation and full occupancy, and neutralize KU-1 17b and 2F5 also

bound less efficiently to HXBc2P3.2, although 2G12 was able to bind both comparably. Si *et al.* [2001]

- IgG1b12: A luciferase-reporter gene-expressing T-cell line was developed to facilitate neutralization and drug-sensitivity assays luciferase and p24 antigen neutralization titer end points were found comparable using NAb from sera from HIV+donors, and MAbs 2F5, 2G12 and IgG1b12. Spenlehauer *et al.* [2001]
- IgG1b12: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 M Abs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001]
- IgG1b12: Twenty HIV clade C isolates from five different countries were susceptible to neutralization by anti-clade B MAbs in a synergistic quadruple combination of mAbs IgG1b12, 2G12, 2F5, and 4E10. Xu et al. [2001]
- IgG1b12: Primary isolates YU2 and ADA are more resistant to IgG1b12 neutralization than HXBc2: 90% Neutralization of HXBc2 is observed with 1.25 ug of IgG1b12, while ADA and YU2 require 2.5 and 5 ug respectively to achieve 50% neutralization, and 90% neutralization could not be achieved with 10 or 20 ug of IgG1b12, respectively. Yang *et al.* [2001]
- IgG1b12: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutralization suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding. York *et al.* [2001]
- IgG1b12: Neutralizing synergy between MAbs 1b12, 2G12 and 2F5 was studied using surface plasmon resonance to determine the binding kinetics for these three MAbs with respect to monomeric and oligomeric env protein gp160 IIIB the 2G12 epitope is highly accessible on both monomeric and oligomeric Envs, 1b12 is highly accessible on monomers but not oligomers, and 2F5 on neither form binding of 2G12 exposes the 2F5 epitope on gp160 oligomers. Zeder-Lutz et al. [2001]
- IgG1b12: b12 recognizes a conformational epitope that overlaps with the CD4 binding site a phage displayed peptide library was used to identify a peptide which bound b12, called B2.1, which competes with b12 in competition assays B2.1 has significant homology to the D loop of gp120: upper case letters indicate residues B2.1 shares with gp120, heRsymFS-DlenrcI one of the goals of defining peptide mimics to the b12 epitope is to develop an immunogen that can stimulate b12-like antibodies, but B2.1 cross-linked to phage and ovalbumin bound IgG1b12 did not elicit cross-reactive gp120 Abs in mice or rabbits. Zwick *et al.* [2001a]
- IgG1b12: This paper primarily concerns 4E10 and Z13, MAbs that both bind proximally to the 2F5 binding site to a conserved epitope, and that neutralize some primary isolates from clades B, C, and E broadly neutralizing MAbs 2F5, IgG1b12, and

4E10 and Z13 fail to neutralize different subsets of viruses. Zwick *et al.* [2001b]

- IgG1b12: Neutralization synergy between anti-HIV NAbs b12, 2G12, 2F5, and 4E10 was studied a classic fixed-ratio method was used, as well as a method where one Ab was fixed at a low neutralization titer and the other was varied using primary isolates, a two-four fold enhancement of neutralization was observed with MAb pairs, and a ten-fold enhancement with a quadruple Ab combination no synergy was observed with any MAb pair in the neutralization of TCLA strain HXB2 whole IgG1b12 and b12 Fab fragments behaved similarly in the neutralization assays there was no evidence for cooperativity of binding between b12 and 2G12 to envelope spikes expressed on the cell surface of TCLA or primary isolates. Zwick *et al.* [2001c]
- IgG1b12: To determine the antigenicity of virus killed by thermal and chemical inactivation, retention of conformationdependent neutralization epitopes was examined, and exposure of CD4BS epitopes was found to be enhanced (MAbs IgG1b12, 205-46-9, and 205-43-1) – binding to 2G12 and 447-52D epitopes was essentially unaltered – the 17b CD4i epitope was also exposed. Grovit-Ferbas *et al.* [2000]
- IgG1b12: SF162 is a neutralization-resistant HIV-1 isolate N-linked glycosylation modifications in the V2 loop of the SF162 gp120 revealed that these sites prevent neutralization by CD4BS MAbs (IgG1b12 and IgGCD4), and protect against neutralization by V3 MAbs (447-D and 391-95D) V2-region glycosylation site mutations did not enhance neutralization resistance to V2 MAbs (G3.4 and G3.136) or CD4i MAbs (17b and 48d) V2 glycosylation site modification allows increased infection of macrophages, probably due to glycosylated forms requiring fewer CCR5 molecules for viral entry. Ly & Stamatatos [2000]
- IgG1b12: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 CD4BS MAbs CD4BS MAbs bound consistently to most isolates of clade D, but poorly to isolates of other clades with the exception of broadly reactive MAb IgG1b12, binding to 22 of 26 isolates tested 8 MAbs were tested for neutralization and MAb IgG1b12 was most potent, with 90% neutralization of 3/5 isolates tested. Nyambi *et al.* [2000]
- IgG1b12: Fab b12 was used six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park *et al.* [2000]
- IgG1b12: rgp120 derived from a R5X4 subtype B virus was used to vaccinate healthy volunteers and the resulting sera were compared with sera from HIV-1 positive subjects and neutralizing MAbs TCLA strains showed enhanced IgG1b12 neutralization sensitivity relative to PBMC-adapted lines IgG1b12 was able to bind, with low affinity, to the rgp120 monomer HIV-1 W61D. Beddows *et al.* [1999]
- IgG1b12: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated

subunits – a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen – SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 – SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 – nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 – anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley *et al.* [1999]

- IgG1b12: Neutralization assays with rsCD4, MAbs, and serum samples from SHIV-infected macaques and HIV-1 infected individuals were used to characterize the antigenic properties of the env glycoprotein of six primary isolate-like or TCLA SHIV variants. IgG1b12 neutralized SHIV strains HXBc2, KU2, 89.6, but not 89.6P and KB9. 89.6 is a dual tropic primary isolate that is not pathogenic in macaques, 89.6P is a highly pathogenic form of 89.6 obtained after passage in macaques, and KB9 is a molecular clone of 89.6P. Neutralization resistance was cell line independent. Crawford *et al.* [1999] (variant cross-recognition or cross-neutralization)
- IgG1b12: The presence of leukocyte function-associated molecule 1 (LFA-1) promotes virus infectivity and hinders neutralization, and anti-LFA-1 MAbs can enhance the neutralizing effect of anti-HIV V3 MAb 447-52D and anti-HIV CD4BS MAb IgG1b12 non-neutralizing anti-HIV CD4BS MAb 654-D did not become neutralizing in the presence of anti-LFA-1 MAbs. Hioe *et al.* [1999]
- IgG1b12: does not inhibit attachment of virus to cells and was used as a control of a study of neutralization by a MAb F58 based micro antibody. Jackson *et al.* [1999]
- IgG1b12: A meeting summary presented results regarding neutralization D. Burton and J. Mascola presented results concerning passive immunization and protection of hu-PBL-SCID mice and macaques, respectively, and both found combinations of MAbs that were able to achieve 99% neutralization in vitro corresponded to efficacy in vivo. Montefiori & Evans [1999]
- IgG1b12: Hu-PBL-SCID mice were infected with HIV-1s JRCSF and SF162 to study the effect of NAbs on an established infection at day 6 post infection, mice were given 50 mg/kg of b12, an amount that would have been protective if given up to 8 hours post-infection, and 100-fold higher than the amount required for 90% neutralization *in vitro* no significant differences in the initial rate of decrease in viral load or the plateau levels of viral RNA between the b12 treated and control mice were seen in most of the Ab treated mice escape mutants were observed with varying patterns of mutations a combination of b12, 2G12 and 2F5 protected 1/3 mice, and an isolate from one of the other two was resistant to neutralization by all three MAbs. Poignard *et al.* [1999]
- IgG1b12: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein (Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer

- CD4BS MAbs 15e, F91 and IgG1b12 bound better to the deleted protein than to wild type. Binley *et al.* [1998]
- IgG1b12: Immunoprecipitation of gp120 and gp160 expressed from a rec Semliki Forest virus by F105 and IgG1b12 indicated that the SFV expressed HIV-1 Env was folded appropriately and SVF-HIV-1 Env vaccine gave the strongest anti-HIV-1 Env response in mice, when compared to an HIV-1 Env DNA vaccine and a rgp160 protein. Brand et al. [1998]
- IgG1b12: Ab from gp120 vaccinated individuals prior to infection, who subsequently became HIV infected, could not achieve 90% neutralization of the primary virus by which the individuals were ultimately infected these viruses were not particularly refractive to neutralization, as determined by their susceptibility to neutralization by MAbs 2G12, IgG1b12, 2F5 and 447-52D. Connor *et al.* [1998]
- IgG1b12: Binds JRSF oligomer with high affinity, as do 205-46-9 and 2G6, but IgG1b12 is neutralizing, the other two are not conclusions of this paper contrast with Parren98 authors propose a model where 205-46-9 and 2G6 may inhibit CD4 binding, but cause a conformational shift which enhances CCR5 binding and thus counteracts the neutralizing effect rank order of CD4BS antibodies oligomer binding is IgG1b12 = 2G6 = 205-46-9 > 205-43-1 = 205-42-15 > 15e = 21h = F91, and the only thing notably distinguishing about neutralizing IgG1b12 is that it depends on residues in V2. Fouts *et al.* [1998]
- IgG1b12: Infection of dendritic cells cultured from CD14+ blood cells or from cadaveric human skin was blocked by neutralizing MAbs IgG1b12, or 2F5 and 2G12 delivered together, but not by control non-neutralizing anti-gp120 MAb 4.8D, indicating that NAbs could interrupt early mucosal transmission events. Frankel *et al.* [1998]
- IgG1b12: anti-C1 region MAb 87-135/9 blocks gp120 interaction with CD4+ cells blocking activity is additive when combined with antibodies which bind in the C4 region of gp120 (F105, 388/389, and b12). Kropelin *et al.* [1998]
- IgG1b12: Enhances binding of Hx10 to CD4 positive or negative HeLa cells, inhibits binding to CD4+ T-cell line A3.01 neutralizes HeLa and A3.01 cell Hx10 infection. Mondor *et al.* [1998]
- IgG1b12: IgG1b12, Fab b12 and 3B3 derived from b12 were all included in this study the rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13 > DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope binding affinity of divalent IgG1b12 is 17-fold greater than monovalent Fab b12. Parren *et al.* [1998a]
- IgG1b12: MAbs 2G12, 2F5 and b12 are broadly neutralizing, as are some human polyconal sera, but this paper describes a set of primary isolates that are resistant to all three MAbs and 2 broadly neutralizing sera results indicate that resistance levels of pediatric isolates might be higher than adult isolates resistance in general did not seem to be conferred by a loss

of binding affinity for gp120 or gp41, rather by a more global perturbation of oligomeric Envelope. Parren *et al.* [1998b]

- IgG1b12: MAbs 654-D100 and IgG1b12 neutralized viruses HIV-BRU and a mutated virus that lacks the V3 loop glycan equally effectively in contrast, sera from guinea pigs immunized with BRU gp120 neutralize viruses more effectively that lack the V3 glycan. Schonning *et al.* [1998]
- IgG1b12: Deleting the V2 loop of neutralization-resistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F deletion of V2, but not V1, diminished neutralization by CD4BS MAb IgG1b12, in contrast to 654.30D and IgGCD4. Stamatatos & Cheng-Mayer [1998]
- IgG1b12: Fab b12 the HIV-1 virus YU2 entry can be enhanced by MAbs binding to the CD4BS, V3 loop, and CD4i epitopes the activation for this enhanced entry state could be conferred on HxB2 by introducing the YU2 V3 loop, or the YU2 V3 and V1/V2 loops a similar effect is observed by subneutralizing concentrations of sCD4 and the effect is dependent of CCR5 Fab fragment b12 also enhances YU2 entry, ruling out Fc interactions or Env cross-linking as a mechanism, while neutralizing HXBc2. Sullivan *et al.* [1998a]
- IgG1b12: Induces Complement-mediated lysis in MN but not primary isolates – primary isolates are refractive to CML. Takefman et al. [1998]
- IgG1b12: MAb was slightly more efficient at neutralization than Fab – inhibits viral binding to cells and viral entry – doesn't affect CD4-independent binding to T-cells. Valenzuela et al. [1998]
- IgG1b12: Summary of the implications of the crystal structure
  of the core of gp120 bound to CD4 and 17b with what is known
  about mutations that reduce NAb binding probable mechanism of neutralization by CD4BS Ab is direct interference with
  CD4 binding IgG1b12 is an unusual CD4BS antibody because it is particularly potent as a neutralizing antibody and it
  is susceptible to changes in the V1-V2 stem loop structure, and
  so it may disrupt an interaction between CD4 and conserved
  amino acids on the V1-V2 stem. Wyatt et al. [1998]
- IgG1b12: Abs that recognize discontinuous epitopes can identify mimotopes from a phage peptide display library IgG1b12 blocks CD4 binding and is the most potent neutralizing Ab many 15 and 21-mer phage inserts were recognized, but it was not possible to derive a consensus common features were a W and at least one acidic residue, and one sequence was found multiple times: NWPRWWEEFVDKHSS, and this peptide could compete with gp120 two short stretches found in the phage peptides might mimic gp120 components of the epitope: positions 382-384, FFY(I), and 423-426 I(FV)I(V)NM. Boots et al. [1997]
- IgG1b12: This is a review that includes a description of IgG1b12, noting approximately equivalent affinities for sgp120 and unprocessed gp160, and somewhat enhanced affinity for the native oligomer on TCLA viruses primary viruses have reduced affinity, but still in the useful range for neutralization there can be complete protection in hu-PBL-SCID mice with Ab even when administered several hours after viral challenge

- competes with sCD4, but unlike other CD4BS antibodies, it is sensitive to mutations in V2. Burton & Montefiori [1997]
- IgG1b12: In a multilab evaluation of monoclonal antibodies, only IgG1b12, 2G12, and 2F5 could neutralize at least half of the 9 primary test isolates at a concentration of < 25 mug per ml for 90% viral inhibition IgG1b12 failed to neutralize only 1/9 primary isolates, although there was some variation between test sites. D'Souza *et al.* [1997]
- IgG1b12: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding IgG1b12 bound monomer, oligomer, and neutralized JRFL. Fouts et al. [1997]
- IgG1b12: b12 was used in its IgG1 form of 14 human MAbs, the most potent neutralizer of SHIV-vpu+, which expressed HIV-1 IIIB env all Ab combinations tested showed synergistic neutralization b12 has a synergistic response with MAbs 694/98-D (anti-V3), 2F5, and 2G12. Li et al. [1997]
- IgG1b12: JRCSF was cultured in the presence of IgG1b12 until a 100-fold resistance to neutralization was selected resistance was due to three changes: V2 substitution D182N and C3 substitution P365L conferred resistance, and V2 D164N was also required for a viable virus IgG1b12 resistant virus remained sensitive to MAbs 2G12 and 2F5. Mo et al. [1997]
- IgG1b12: Review: MAbs 2F5, 2G12 and IgG1b12 have potential for use in combination with CD4-IgG2 as an immunotherapeutic or immunoprophylactic homologous MAbs to these are rare in humans and vaccine strategies should consider including constructs that may enhance exposure of these MAbs' epitopes. Moore & Trkola [1997]
- IgG1b12: Complete protection against HIV-1 infection was achieved in hu-PBL-SCID mice by passive immunization with physiologically relevant doses pharmacokinetics showed serum half-life of 30.2 +/- 1.3 hours for Fab b12 and 7.4 +/- 0.7 days for IgG1 b12 in mice, but IgG1 half-lives in human are generally between 21-23 days. Parren *et al.* [1995]; Parren & Burton [1997] (immunoprophylaxis)
- IgG1b12: In this review, the technique and potential application of Fab expression and selection in phage display libraries, and subsequent production of IgG molecules is discussed b12 is exceptionally potent at neutralization and can successfully neutralize most B clade primary isolates, and many isolates from other subtypes as well 3B3 was derived from b12 by selection for higher affinity using the CDR walking strategy 3B3 has 8-fold enhancement of binding, a linear correlation was found between neutralization and affinity, and 3B3 can neutralize strains b12 cannot. Parren & Burton [1997]
- IgG1b12: Fab b12 is unusual in that it binds to gp140 and monomeric gp120 with similar affinities, and with a higher affinity to the native oligomer—authors propose this antibody may be exceptional because it binds the virus rather than viral debris—IgG1b12 can protect against infection prior to or shortly after challenge of hu-PBL-SCID mice with TCLA strains and primary strains, but the serum concentrations required *in vivo* were higher than for *in vitro* neutralization. Parren et al. [1997b,a]
- IgG1b12: Inhibited some SI- and NSI-env chimeric viruses but enhanced one NSI-env chimeric virus 3 fold. Schutten *et al.* [1997]

- IgG1b12: Viral binding inhibition by IgG1b12 strongly correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5). Ugolini et al. [1997]
- IgG1b12: Major deletions in C1 and C5 and deletions of the V1V2 and V3 loops do not diminish binding. Wyatt *et al.* [1997]
- IgG1b12: Saturation mutagenesis of the complementaritydetermining region and optimization strategies were used to create very high affinity versions of this Fab – increased affinity was dominated by a slowing of the off rate. Yang et al. [1997c]
- IgG1b12: 35 primary isolates were tested and all were neutralized by IgG1b12 (including 4, UG270, RW92/026, ZB20, and 301727 which been had reported as not neutralized by IgG1b12 Trkola *et al.* [1995]) IgG1b12 could neutralize even when added after the virus to the culture selection for 400-fold increased affinity did not enhance neutralization by antibody IgG1b12 was more potent with greater breadth than MAb 2F5 Kessler II *et al.* [1997]. Kessler II *et al.* [1997]; Trkola *et al.* [1995]
- IgG1b12: Potent neutralizing *ex vivo* of virus taken directly from plasma of HIV-1 infected individuals little correlation between neutralization sensitivity of passaged virus and plasma derived virus more effective than MAb 19b. Gauduin *et al.* [1996]
- IgG1b12: Review: Unique among anti-CD4BS MAbs in terms of being potent against both lab adapted virus and primary isolates one of three MAbs (IgG1b12, 2G12, and 2F5) generally accepted as having significant potency against primary isolates. Poignard *et al.* [1996b]
- IgG1b12: Anti-CD4BS MAbs 15e, 21h, and IgG1b12 did not cause gp120 dissociation from virus, or exposure of the gp41 epitope of MAb 50-69, in contrast to CD4i MAb 48d and anti-V3 neutralizing MAbs. Poignard *et al.* [1996a]
- IgG1b12: Review: Only four epitopes have been described which can stimulate a useful neutralizing response to a broad spectrum of primary isolates, represented by the binding sites of MAbs: 447-52-D, 2G12, Fab b12, and 2F5. Sattentau [1996]
- IgG1b12: Neutralizes JR-FL inhibits gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola *et al.* [1996a]
- IgG1b12: Because Fab b12 shows reduction in binding when the V2 loop is deleted and when aa 183/184 PI/SG substitutions are made competition studies were done with Fab L78 and anti-V2 MAbs SC258 and 684-238 and they do not compete with IgG1b12. Ditzel *et al.* [1995]
- IgG1b12: Called BM12 broad cross-clade neutralization of primary isolates additive neutralization in combination with MAb 2F5. Kessler *et al.* [1995] (antibody interactions)
- IgG1b12: Anti-CD4 binding site MAb very potent neutralization of a number of primary isolates. Moore *et al.* [1995a]
- IgG1b12: Review: unusual properties for anti-CD4 BS MAb: sensitive to V2 substitutions, preferential recognition of the oligomer on the cell surface. Moore & Ho [1995]
- IgG1b12: Formalin inactivation of virus at 0.1% formalin for 10 hours at 4 degrees was optimal for inactivation of virus while maintaining epitope integrity. Sattentau *et al.* [1995]
- IgG1b12: Fab b12 showed potent neutralization of T-cell-line-adapted strains, but much reduced neutralization of 3 primary isolates 2 of the 3 primary isolates also had reduced binding

- affinity, but the third was as efficiently immunoprecipitated as HXBc2. Sullivan *et al.* [1995]
- IgG1b12: Could potently neutralize primary isolates from within clade B, but showed a slight reduction in efficacy outside of clade B. Trkola *et al.* [1995]
- IgG1b12: Very potent neutralization, of primary and lab strains, at concentrations that could be achieved by passive immunization reduced binding with A,C, and D clade viruses relative to B clade, poor reactivity with E clade isolates that were refractive to neutralization by sera from HIV-1 + donors could be neutralized by IgG1 b12. Burton *et al.* [1994] (variant cross-recognition or cross-neutralization)
- IgG1b12: Cross-reactive with some gp120s, (but not all), from clades A-D – not reactive with gp120 from clades E or F. Moore *et al.* [1994b] (variant cross-recognition or crossneutralization)
- IgG1b12: Anti-CD4 binding site Fab, potent neutralizing activity, greater affinity for a subpopulation of gp120 molecules suggested to be in a mature confirmation mutations in gp120 that abrogate binding: 368 D/R or D/T, 370 E/R, and 477 D/V, of clone HXBc2 of LAI sensitive to V1 and V2 substitutions. Roben *et al.* [1994] (antibody binding site definition and exposure)
- IgG1b12: The original Fab fragment was derived from a combinatorial phage library from bone marrow of an HIV-1 positive individual who had been asymptomatic for six years. Burton *et al.* [1991] (antibody generation)

**No.** 1087

MAb ID IgGCD4 (IgG-CD4)

**HXB2 Location** Env **Author Location** gp120

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human (IgG)

Ab Type gp120 CD4BS

References Srivastava et al. 2002; Ly & Stamatatos 2000; Stamatatos & Cheng-Mayer 1998; Capon et al. 1989

- IgGCD4: Oligomeric gp140 (o-gp140) derived from R5 primary isolate US4 was characterized for use as a vaccine reagent

   antigen capture ELISA was used to compare the antigenicity of gp120 and o-gp140 using a panel of well characterized MAbs Abs directed against the CD4 binding site (IgGCD4 and IgG1b12) reacted slightly more strongly with the gp120 monomer than with the oligomer, as did sCD4. Srivastava et al. [2002]
- IgGCD4: SF162 is a neutralization-resistant HIV-1 isolate N-linked glycosylation modifications in the V2 loop of the SF162 gp120 revealed that these sites prevent neutralization by CD4BS MAbs (IgG1b12 and IgGCD4), and protect against neutralization by V3 MAbs (447-D and 391-95D) V2-region glycosylation site mutations did not alter neutralization resistance to V2 MAbs (G3.4 and G3.136) or CD4i MAbs (17b and 48d) V2 glycosylation site modification allows infection of macrophages, probably due to glycosylated forms requiring fewer CCR5 molecules for viral entry. Ly & Stamatatos [2000]

• IgGCD4: Deleting the V2 loop of neutralization-resistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F – deletion of V2 but not V1 slightly enhanced neutralization by CD4BS MAb IgGCD4. Stamatatos & Cheng-Mayer [1998]

• IgGCD4: An antibody-like immunoadhesins molecule was constructed incorporating the gp120-binding domain of CD4. Capon *et al.* [1989]

No. 1088

MAb ID L28

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type gp120 CD4BS

**References** Gorny & Zolla-Pazner 2004; Ditzel *et al.* 1995 **Keywords** antibody binding site definition and exposure, antibody sequence, variable domain, review

- L28: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- L28: Substitutions at 257 T/R, 368 D/R, 370 E/R and 370 E/Q, 475 M/S 102 E/L and 463 N/D reduce binding binding was enhanced by removal of the V3 loop and by substitutions 45 W/S, 298 R/G, 381 E/P, 382 F/L, 420 I/R, 435 Y/H or Y/R binding is sensitive to deglycosylation heavy and light chain variable region sequence is available. Ditzel *et al.* [1995] (antibody binding site definition and exposure, antibody sequence, variable domain)

**No.** 1089

MAb ID L33

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**Ab Type** gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Zwick et al.

2003; Ditzel et al. 1995

**Keywords** antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain, review

- L33: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- L33: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs

directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick *et al.* [2003] (antibody interactions)

 L33: binding is sensitive to deglycosylation – heavy and light chain variable region sequence is available. Ditzel *et al.* [1995] (antibody binding site definition and exposure, antibody sequence, variable domain)

**No.** 1090

MAb ID L41

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

**Neutralizing** L

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**Ab Type** gp120 CD4BS

**References** Gorny & Zolla-Pazner 2004; Ditzel *et al.* 1995 **Keywords** antibody binding site definition and exposure, antibody sequence, variable domain, review

- L41: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- L41: Substitutions at 133 D/R, 256 S/Y, 257 T/R, 368 D/R or D/T, 370 E/Q or E/R, 384 Y/E, and 421 K/L reduce binding paradoxically, this Fab was retrieved from the library after masking with known anti-CD4BS MAbs binding is sensitive to deglycosylation heavy and light chain variable region sequence is available. Ditzel *et al.* [1995] (antibody binding site definition and exposure, antibody sequence, variable domain)

**No.** 1091

MAb ID L42

HXB2 Location Env

**Author Location** gp120

**Epitope** 

**Neutralizing** L

Immunogen HIV-1 infection

**Species (Isotype)** human ( $IgG1\kappa$ )

**Ab Type** gp120 CD4BS

**References** Gorny & Zolla-Pazner 2004; Ditzel *et al.* 1995 **Keywords** antibody binding site definition and exposure, review

- L42: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- L42: Substitutions at 257 T/R, 368 D/R, 370 E/R, 266 A/E and 477 D/V reduce binding binding was significantly enhanced by 381 E/P and 382 F/L binding is sensitive to deglycosylation heavy and light chain variable region sequence is available. Ditzel *et al.* [1995] (antibody binding site definition and exposure)

**No.** 1092

MAb ID L52
HXB2 Location Env
Author Location gp120
Epitope
Neutralizing L
Immunogen HIV-1 infection
Species (Isotype) human (IgG1κ)

**Ab Type** gp120 CD4BS **References** Gorny & Zolla-Pazner 2004; Ditzel *et al.* 1995 **Keywords** antibody binding site definition and exposure,

antibody sequence, variable domain, review

 L52: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)

• L52: Binding is sensitive to deglycosylation – heavy and light chain variable region sequence is available. Ditzel *et al.* [1995] (antibody binding site definition and exposure, antibody sequence, variable domain)

No. 1093

MAb ID L72

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing Immunogen

Species (Isotype) mouse

**Ab Type** gp120 CD4BS

Research Contact Dr. Hariharam, IDEC Pharmaceuticals Corp La Jolla, CA

References Ditzel et al. 1997

• L72: Used to bind gp120 to solid phase to select MAbs from a phage selection library. Ditzel *et al.* [1997]

**No.** 1094

MAb ID M12

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

**Research Contact** P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- M12: There is a p15 gag specific MAb also named M12.
- M12: A comparison of 25 gp120 specific, conformation dependent MAbs was done M12 is part of a group of MAbs labeled A1 all A1 MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, blocked CD4 binding, were sensitive to mutations in gp120 positions 368 and 370 that directly contact CD4 50% neutralization of NL4-3 was achieved with 21 ug/ml of M12. Sugiura *et al.* [1999]
- M12: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

**No.** 1095

MAb ID M13

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

• L52: This review summarizes MAbs directed to HIV-1 Env. **Research Contact** P. Earl, National Institute of Allergy and In-There are 51 CD4BS MAbs and Fabs in the database; most, like fectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- M13: A comparison of 25 gp120 specific, conformation dependent MAbs was done M13 is part of a group of MAbs labeled A1 all A1 MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, blocked CD4 binding, were sensitive to mutations in gp120 positions 368 and 370 that directly contact CD4 50% neutralization of NL4-3 was achieved with 35 ug/ml of M13. Sugiura et al. [1999]
- M13: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1096

MAb ID M6

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

**Species** (**Isotype**) mouse (IgG)

Ab Type gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- M6: A comparison of 25 gp120 specific, conformation dependent MAbs was done M6 is part of a group of MAbs labeled A1 all A1 MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, blocked CD4 binding, were sensitive to mutations in gp120 positions 368 and 370 that directly contact CD4. Sugiura et al. [1999]
- M6: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1097

MAb ID MAG 116

**HXB2 Location** Env

Author Location gp120

Epitope

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 CD4BS

Research Contact C. Y. Kang, IDEC Inc

References Kang et al. 1994

MAG 116: Amino acid substitutions that reduce binding 10 fold: 256 S/Y, 257 T/R, 368 D/R or T, 370 E/R or Q, 384 Y/E, 421 K/L – neutralizes MN, IIIB and RF. Kang et al. [1994]

No. 1098

MAb ID MAG 12B

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 CD4BS

Research Contact C. Y. Kang, IDEC Inc

References Kang et al. 1994

MAG 12B: Amino acid substitutions that reduce binding 10 fold: 257 T/R, 368 D/R or T, 370 E/R or Q, 384 Y/E, 477 D/V – weak neutralization of IIIB. Kang et al. [1994]

No. 1099

MAb ID MAG 29B

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 CD4BS

Research Contact C. Y. Kang, IDEC Inc

References Kang et al. 1994

MAG 29B: Amino acid substitutions that reduce binding 10 fold: 257 T/R, 368 D/R or T, 370 E/R or Q, 384 Y/E, 386 N/Q, 421 K/L – weak neutralization of IIIB. Kang *et al.* [1994]

**No.** 1100

MAb ID MAG 3B

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 CD4BS

Research Contact C. Y. Kang, IDEC Inc

References Kang et al. 1994

MAG 3B: Amino acid substitutions that reduce binding 10 fold:
 256 S/Y, 257 T/R or A or G, 262 N/T, 368 D/R or T, 370 E/R or Q, 381 E/P, 384 Y/E, 421 K/L, 475 M/S, 477 D/V. Kang et al. [1994]

**No.** 1101

**MAb ID** MAG 55 (#55)

HXB2 Location Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 CD4BS

Research Contact C. Y. Kang, IDEC Inc

**References** Koefoed *et al.* 2005; Moore & Sodroski 1996; Kang *et al.* 1994

Keywords antibody binding site definition and exposure

- MAG 55: IgG antibody phage display libraries were created from HIV-1 + individuals after pre-selection of PBMC with gp120, as an alternative to using bone marrow for generating libraries. MAG 55 was among a set of Abs used for competition studies to define the binding sites of the newly isolated MAbs, representing a MAb with a CD4BS epitope. Koefoed *et al.* [2005] (antibody binding site definition and exposure)
  - MAG 55: Called #55 binding reciprocally inhibited by other anti-CD4 binding site MAbs, and by some C1-C5 MAbs – binding enhanced by anti-V3 MAb 110.5 and anti-V2 MAbs G3-136 and G3-4 – enhances binding of many anti-V3 and -V2 MAbs. Moore & Sodroski [1996]
- MAG 55: Amino acid substitutions that reduce binding 10 fold: 256 S/Y, 257 T/R, 368 D/R or T, 370 E/R or Q, 384 Y/E, 421 K/L, 470 P/L, 475 M/S, 477 D/V – neutralizes MN, IIIB and RF. Kang et al. [1994]

**No.** 1102

**MAb ID** MAG 72 (L72)

HXB2 Location Env

**Author Location** gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 CD4BS

Research Contact C. Y. Kang or Dr. Hariharam, IDEC Pharmaceuticals Corp, La Jolla, CA

References Ditzel et al. 1997; Kang et al. 1994

- MAG 72: Called L72 used to bind gp120 to solid phase to select MAbs from a phage selection library. Ditzel *et al.* [1997]
- MAG 72: Amino acid substitutions that reduce binding 10 fold: 257 T/R or A or G, 262 N/T, 368 D/R or T, 370 E/R or Q, 384 Y/E, 421 K/L, 477 D/V – neutralizes MN, IIIB and RF. Kang et al. [1994]

**No.** 1103

MAb ID MAG 86

**HXB2 Location** Env

Author Location gp120

Epitope

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 CD4BS

Research Contact C. Y. Kang, IDEC Inc

References Kang et al. 1994

 MAG 86: Amino acid substitutions that reduce binding 10 fold: 256 S/Y, 257 T/R, 368 D/R or T, 370 E/R or Q, 384 Y/E, 421 K/L, 470 P/L, 477 D/V – neutralizes MN, IIIB and RF. Kang et al. [1994]

No. 1104

MAb ID MAG 96

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 CD4BS

Research Contact C. Y. Kang, IDEC Inc

References Koefoed et al. 2005; Kang et al. 1994

**Keywords** antibody binding site definition and exposure

- MAG 96: IgG antibody phage display libraries were created from HIV-1 + individuals after pre-selection of PBMC with gp120, as an alternative to using bone marrow for generating libraries. MAG 96 was among a set of Abs used for competition studies to define the binding sites of the newly isolated MAbs, representing a MAb with a CD4BS epitope. Koefoed *et al.* [2005] (antibody binding site definition and exposure)
- MAG 96: Amino acid substitutions that reduce binding 10 fold: 256 S/Y, 257 T/R, 368 D/R or T, 370 E/R – weak neutralization of IIIB. Kang et al. [1994]

No. 1105

MAb ID MTW61D

**HXB2 Location** Env

Author Location gp120 (W61D)

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human

**Ab Type** gp120 CD4BS

**References** Gorny & Zolla-Pazner 2004; Fouts *et al.* 1998; Sullivan *et al.* 1998a

**Keywords** enhancing activity, review

- MTW61D: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- MTW61D the HIV-1 virus YU2 entry can be enhanced by MAbs binding to the CD4BS, V3 loop, and CD4i epitopes the activation for this enhanced entry state could be conferred on HxB2 by introducing the YU2 V3 loop, or the YU2 V3 and V1/V2 loops a similar effect is observed by sub-neutralizing concentrations of sCD4 and the effect is dependent of CCR5 Fab fragment MTW61D also enhances YU2 entry, ruling out Fc interactions or Env cross-linking as a mechanism, while neutralizing HXBc2 MTW61D was obtained by panning libraries derived from bone marrow from a long term asymptomatic

donor against gp120 from primary isolate W61D. Sullivan *et al.* [1998a] (**enhancing activity**)

**No.** 1106

MAb ID S1-1

**HXB2** Location Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 CD4BS

**References** Gorny & Zolla-Pazner 2004; Wisnewski *et al.* 1996; Moran *et al.* 1993; Lake *et al.* 1992

**Keywords** antibody binding site definition and exposure, antibody sequence, variable domain, comple-

ment, enhancing activity, review

• S1-1: This review summarizes MAbs directed to HIV-1 Env.
There are 51 CD4BS MAbs and Fabs in the database; most, like

- this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (**review**)
   S1-1: S1-1 is V H1 V-region heavy chain usage was examined and a bias of enhanced V H1 and V H4, and reduced V H3,
- and a bias of enhanced V H1 and V H4, and reduced V H3, was noted among HIV infected individuals. Wisnewski *et al.* [1996] (antibody sequence, variable domain)
- S1-1: Heavy (V HI) and light (V lambdaIII) chain sequenced

   no enhancing activity similar germline sequence to MAb
   86, but very different activity. Moran *et al.* [1993] (enhancing activity, antibody sequence, variable domain)
- S1-1: Neutralizes IIIB and MN without complement, and neutralizes RF and a clinical isolate with complement binds to native but not denatured gp120 inhibits sCD4-gp120 binding. Lake *et al.* [1992] (antibody binding site definition and exposure, complement)

**No.** 1107

MAb ID T13

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope

Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type gp120 CD4BS

• MTW61D: This review summarizes MAbs directed to HIV-1 **Research Contact** P. Earl, National Institute of Allergy and In-Env. There are 51 CD4BS MAbs and Fabs in the database; fectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- T13: A comparison of 25 gp120 specific, conformation dependent MAbs was done T13 is one of three MAbs labeled group Cb, that was type-specific for BH8 T13 fully blocked CD4 binding, and the deletion of the V3 loop enhanced binding 10-fold. Sugiura *et al.* [1999]
- T13: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1108 **MAb ID** T49

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope Neutralizing** no **Immunogen** vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

**Ab Type** gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- T49: A comparison of 25 gp120 specific, conformation dependent MAbs was done T49 is one of three MAbs labeled group Cb, that was type-specific for BH8 T49 fully blocked CD4 binding, and the deletion of the V3 loop enhanced binding 10-fold. Sugiura *et al.* [1999]
- T49: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

No. 1109

MAb ID T56

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope
Neutralizing no
Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

**Species (Isotype)** mouse (IgG) **Ab Type** gp120 CD4BS

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- T56: A comparison of 25 gp120 specific, conformation dependent MAbs was done T56 is one of three MAbs labeled group Cb, that was type-specific for BH8 T56 fully blocked CD4 binding, and the deletion of the V3 loop enhanced binding 10-fold. Sugiura *et al.* [1999]
- T56: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

**No.** 1110

MAb ID TH9

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type gp120 CD4BS

Research Contact Michael Fung, Tanox Biosystem, USA

References Gorny & Zolla-Pazner 2004; Yang et al. 1998;

D'Souza et al. 1995

**Keywords** assay development, review, subtype comparisons, variant cross-recognition or cross-

neutralization

- TH9: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this MAb, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- TH9: A neutralization assay was developed based on heminested PCR amplification of the LTR (HNPCR) LTR-HNPCR consistently revealed HIV DNA and was shown to be a rapid, specific and reliable neutralization assay based on tests with 6 MAbs and 5 isolates. Yang et al. [1998] (assay development)
- TH9: Found to neutralize MN, but not JRCSF, two B subtype primary isolates, or a D subtype primary isolate, by most labs in a multi-laboratory study involving 11 labs. D'Souza *et al.* [1995] (variant cross-recognition or cross-neutralization, subtype comparisons)

No. 1111

MAb ID anti-CD4BS summary

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen

Species (Isotype)

Ab Type gp120 CD4BS

References Moore & Sodroski 1996; Thali et al. 1993

- Anti-CD4 binding site antibodies (CD4BS) competitively inhibit CD4 binding to monomeric gp120, and they differ in precise dependence on gp120 residues, but generally require Asp-368 and Glu-370. Moore & Sodroski [1996]
- Shared components of MAb epitopes and the discontinuous CD4 binding regions included Thr 257, Asp 368, Glu 370, Lys 421 through Trp 427 and Asp 457. Thali et al. [1993]

**No.** 1112

MAb ID b11

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 CD4BS

**References** Gorny & Zolla-Pazner 2004; Parren *et al.* 1998a

**Keywords** binding affinity, review

- b11: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- b11: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13 > DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren *et al.* [1998a] (**binding affinity**)

No. 1113
MAb ID b13
HXB2 Location Env
Author Location gp120
Epitope
Neutralizing
Immunogen
Species (Isotype) human

**Ab Type** gp120 CD4BS

**References** Gorny & Zolla-Pazner 2004; Parren & Burton 1997; Parren *et al.* 1998a; Parren *et al.* 1995

Keywords binding affinity, immunoprophylaxis, review

- b13: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- b13: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13 > DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren *et al.* [1998a] (**binding affinity**)
- b13: Fab b13 was used as a control in a hu-PBL SCID mouse study animals were protected from HIV-1 SF2 infection by IgG1b12, somewhat by Fab b12, but not by b13. Parren *et al.* [1995]; Parren & Burton [1997] (immunoprophylaxis)

No. 1114
MAb ID b14
HXB2 Location Env
Author Location gp120
Epitope
Neutralizing
Immunogen
Species (Isotype) human

Ab Type gp120 CD4BS

References Gorny & Zolla-Pazner 2004; Parren et al. 1998a

Keywords binding affinity, review

- b14: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- b14: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13 > DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren *et al.* [1998a] (**binding affinity**)

**No.** 1115 **MAb ID** b3

HXB2 Location Env
Author Location gp120
Epitope
Neutralizing
Immunogen
Species (Isotype) human
Ab Type gp120 CD4BS

References Selvarajah et al. 2005; Pantophlet et al. 2004; Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2003: Pantophlet et al.

2003b; Zwick *et al.* 2003; Pantophlet *et al.* 2003a; Parren *et al.* 1998a; Parren *et al.* 1997b

**Keywords** antibody binding site definition and exposure, antibody interactions, binding affinity, review, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-neutralization

- b3: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. CD4BS MAbs except Fab b12 (b6, b3, F105) did not bind to either GDMR or mCHO. Selvarajah *et al.* [2005] (vaccine antigen design, vaccine-specific epitope characteristics)
- b3: This review summarizes MAbs directed to HIV-1 Env. There are 51 CD4BS MAbs and Fabs in the database; most, like this Fab, neutralize TCLA strains only. Gorny & Zolla-Pazner [2004] (review)
- b3: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including b3. Pantophlet *et al.* [2004] (vaccine antigen design)
- b3: Alanine scanning mutagenesis was used to compare substitutions that affected anti-CD4BS NAb b12 binding to those that affect binding of sCD4 and two non-neutralizing anti-CD4BS Abs b3 and b6 while the epitope maps overlapped, there were some differences observed binding of CD4 was never enhanced, indicating it had evolved to be optimal rec gp120s were engineered to contain combinations of Alanine substitutions that enhanced b12 binding, and while binding of b12 to these gp120 monomers was generally maintained or increased, binding by five non-neutralizing anti-CD4bs MAbs (b3, b6, F105, 15e, and F91) was reduced or completely abolished 2G12 binding was largely unperturbed, indicating these proteins were not grossly misfolded. Pantophlet *et al.* [2003a] (antibody binding site definition and exposure)
- b3: A gp120 molecule was design to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of

non-neutralizing MAbs were combined with additional Nlinked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)

- b3: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick et al. [2003] (antibody interactions)
- b3: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13 >DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs - authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a] (binding affinity)
- b3: Neutralizes TCLA strains, but not primary isolates. Parren et al. [1997b] (variant cross-recognition or crossneutralization)

**No.** 1116

MAb ID b6

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

**Immunogen** 

Species (Isotype) human

Ab Type gp120 CD4BS

Research Contact Dennis Burton, Scripps, San Diego, CA, USA

References Selvarajah et al. 2005; Pantophlet et al. 2004; Binley et al. 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Pantophlet et al. 2003a; Poignard et al. 2003; Kwong et al. 2002; Parren et al. 1998a; Parren et al. 1997b

**Keywords** antibody binding site definition and exposure, antibody interactions, subtype comparisons, vaccine antigen design, vaccine-specific epitope characteristics, variant cross-recognition or cross-neutralization

• b6: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. CD4BS MAbs except Fab b12 (b6, b3, F105) did not bind to either GDMR

or mCHO. Selvarajah et al. [2005] (vaccine antigen design, vaccine-specific epitope characteristics)

- b6: 93 viruses from different clades were tested for their neutralization cross-reactivity using a panel of HIV antibodies. b6 was included as an example of a CD4BS antibody that is not strongly neutralizing, and it only was able to neutralize a few highly sensitive primary viruses and T-cell adapted viral strains that were B clade. Steric restrictions probably block its binding site in most isolates. Binley et al. [2004] (variant cross-recognition or cross-neutralization, subtype compar-
- b6: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including b6. Pantophlet et al. [2004] (vaccine antigen design)
- b6: Alanine scanning mutagenesis was used to compare substitutions that affected anti-CD4BS NAb b12 binding to those that affect binding of sCD4 and two non-neutralizing anti-CD4BS Abs b3 and b6 - while the epitope maps overlapped, there were some differences observed – binding of CD4 was never enhanced, indicating it had evolved to be optimal - rec gp120s were engineered to contain combinations of Alanine substitutions that enhanced b12 binding, and while binding of b12 to these gp120 monomers was generally maintained or increased, binding by five non-neutralizing anti-CD4bs MAbs (b3, b6, F105, 15e, and F91) was reduced or completely abolished -2G12 binding was largely unperturbed, indicating these proteins were not grossly misfolded. Pantophlet et al. [2003a]
- b6: A gp120 molecule was design to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional Nlinked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)
- b6: Virion capture assays are not a good predictor of neutralization, and the presentation of epitopes using this assay seems to be different from that of functional Envelope spikes on primary isolates – F105 and b6 could efficiently block the b12-mediated capture of infectious virions in a virus capture, but did not inhibit b12 neutralization – while b12 was potent at neutralizing the three primary virions JR-CSF, ADA, and 89.6, the Abs F105, 19b, and Fab b6 were overall very poor neutralizers. Poignard et al. [2003]
- b6: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics

of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the CD4BS MAbs used. Zwick *et al.* [2003] (antibody interactions)

- b6: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar. Enthalpy and entropy changes were divergent, but compensated. Not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs (17b, 48d, 1.5e, b6, F105 and F91) had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, but the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. The high values suggest surface burial or protein folding an ordering of amino acids. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong et al. [2002] (antibody binding site definition and exposure)
- b6: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13 > DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren *et al.* [1998a]
- b6: Neutralizes TCLA strains, but not primary isolates. Parren et al. [1997b]

No. 1117 MAb ID polyclonal HXB2 Location Env Author Location gp120

Epitope Neutralizing no Immunogen vaccine

Vector/Type: protein, virus-like particle (VLP) Strain: B clade LAI HIV component: CD4BS, Gag, V3

Species (Isotype) mouse

**Ab Type** gp120 CD4BS **References** Truong *et al.* 1996

Antibodies raised against recombinant anti-p55 virus-like particles with the p24 region 196-226 deleted, bearing inserts of either the V3 or the CD4BS regions of gp120 were studied – no neutralizing responses, weak Env and strong Gag responses were elicited – the major homology region (MHR) and proximal sequences was found to be required for capsid assembly. Truong et al. [1996]

**No.** 1118

MAb ID
HXB2 Location Env
Author Location gp120
Epitope

Neutralizing yes Immunogen

Species (Isotype) human

**Ab Type** gp120 CD4BS, gp120 CD4i, gp120 V2, gp120 V3

References Moore et al. 2001

• Moore and colleagues review structural aspects of gp120 and how they relate to antigenic domains, and review the data concerning the lack of a clear relationship between genetic subtype and serotype – they suggest the primary goal in vaccine efforts should be to design an immunogen that can be shown to elicit neutralizing antibodies against a significant proportion of primary isolates – assay artifacts that can result in confused interpretations are also discussed, such as Ab binding to defective spikes, which does not affect HIV-1 infectivity, but can dominant an assay signal. Moore et al. [2001]

No. 1119

**MAb ID** 17b (1.7b, sCD4-17b)

HXB2 Location Env Author Location gp120 Epitope

Neutralizing L P (weak)
Immunogen HIV-1 infection

Species (Isotype) human

**Ab Type** gp120 CD4i, gp120 CCR5BS

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leans, LA, USA

References Selvarajah et al. 2005; Martín-García et al. 2005; Lusso et al. 2005; Koefoed et al. 2005; Haynes et al. 2005; Pinter et al. 2004; Pantophlet et al. 2004; Nabatov et al. 2004; Mc-Caffrey et al. 2004; Ling et al. 2004; Liao et al. 2004; Biorn et al. 2004; Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Zhu et al. 2003; Ohagen et al. 2003; Xiang et al. 2003; Labrijn et al. 2003; Enshell-Seijffers et al. 2003; Dey et al. 2003; Cavacini et al. 2003; Binley et al. 2003; He et al. 2003; Ling et al. 2002; Finnegan et al. 2002; Cavacini et al. 2002; Arthos et al. 2002; Zhang et al. 2002; Basmaciogullari et al. 2002; Grundner et al. 2002; Edwards et al. 2002; Xiang et al. 2002a; Xiang et al. 2002b; Dowd et al. 2002; Yang et al. 2002; Schulke et al. 2002; Golding et al. 2002b; Srivastava et al. 2002; Kwong et al. 2002; Finnegan et al. 2001; Poignard et al. 2001; Zhang et al. 2001a; York et al. 2001; Kolchinsky et al. 2001; Si et al. 2001; Rizzuto & Sodroski 2000; Yang et al. 2000; Stamatatos et al. 2000; Salzwedel et al. 2000; Park et al. 2000; Ly & Stamatatos 2000; Grovit-Ferbas et al. 2000; Binley et al. 1999; Hoffman et al. 1999; Oscherwitz et al. 1999a; Stamatatos &

Cheng-Mayer 1998; Binley et al. 1998; Sullivan et al. 1998a; Sullivan et al. 1998; Rizzuto et al. 1998; Moore & Binley 1998; Wyatt et al. 1998; Kwong et al. 1998; Parren et al. 1997b; Wyatt et al. 1997; Cao et al. 1997b; Ditzel et al. 1997; Weinberg et al. 1997; Li et al. 1997; Fouts et al. 1997; Binley et al. 1997a; Trkola et al. 1996a; Wu et al. 1996; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; Wyatt et al. 1995; Beretta & Dalgleish 1994; Thali et al. 1994; Moore et al. 1993c; Thali et al. 1993

**Keywords** vaccine antigen design, vaccine-specific epitope characteristics

- 17b: 48d and 17b have similar epitopes, and the pair are unique among human and rodent MAbs.
- 17b: NIH AIDS Research and Reference Reagent Program: 4091.
- 17b: Called 1.7B. Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 1.7B has no indication of polyspecific autoreactivity. Haynes et al. [2005]
- 17b: IgG antibody phage display libraries were created from HIV-1 + individuals after pre-selection of PBMC with gp120, as an alternative to using bone marrow for generating libraries.
   17b was among a set of Abs used for competition studies to define the binding sites of the newly isolated MAbs, representing a MAb with a CD4i epitope. Koefoed *et al.* [2005]
- 17b: The epitope for the MAb D19 is conserved and embedded in V3. D19 is unique in that for R5 viruses, it was cryptic and did not bind without exposure to sCD4, and for X4 and R5X4 isolates it was constitutively exposed. D19b is unque among CD4i antibodies in that it binds to the V3 loop. CD4i MAbs 17b and 48d were used as controls for CD4i characterization; in contrast to D19, other CD4i MAbs bind to the conserved bridging sheet and do not differentiate between R5 and X4 using strains. 17b, like D19, was able to neutralize the BaL isolate only in combination with sCD4. Lusso *et al.* [2005]
- 17b: The HIV-1 Bori-15 variant was adapted from the Bori isolate for replication microglial cells. Bori-15 had increased replication in microglial cells and a robust syncytium-forming phenotype, ability to use low levels of CD4 for infection, and increased sensitivity to neutralization by sCD4 and 17b. Four amino acid changes in gp120 V1-V2 were responsible for this change. Protein functionality and integrity of soluble, monomeric gp120-molecules derived from parental HIV-1 Bori and microglia-adapted HIV-1 Bori-15 was assessed in ELISA binding assays using F105, IgG1b12, 17b and 48d, 2G12 and 447-52D. Association rates of sCD4 and 17b were not changed, but dissociation rates were 3-fold slower for sCD4 and 14-fold slower for 17b. Martín-García et al. [2005]
- 17b: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did

not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. CD4i MAbs (48d, 17b) did not bind to either GDMR or mCHO even with sCD4. Selvarajah *et al.* [2005] (vaccine antigen design, vaccine-specific epitope characteristics)

- 17b: The peptide 12p1 (RINNIPWSEAMM) inhibits direct binding of YU2 gp120 or Env trimer to CD4, CCR5 and MAb 17b in a concentration-dependent allosteric manner. 12p1 is thought to bind to unbound gp120 near the CD4 binding site, with a 1:1 stoichiometry. 12p1 also inhibited MAb F105 binding presumably because F105 favors an unactivated conformation, but not MAbs 2G12 or b12. The 1:1 stoichiometry, the fact that the peptide binding site is accessible on the trimer, the non-CD4 like aspect of the binding, and an ability to inhibit viral infection in cell cultures make it a promising lead for therapeutic design. Biorn *et al.* [2004]
- 17b: This review summarizes MAbs directed to HIV-1 Env. There are six CD4 inducible MAbs and Fabs in the database. The MAb forms neutralize TCLA strains only, but the smaller Fabs and scFv fragments can neutralize primary isolates. Gorny & Zolla-Pazner [2004]
- 17b: A32-rgp120 complexes opened up the CCR5 co-receptor binding site, but did not induce neutralizing antibodies with greater breadth among B subtype isolates than did uncomplexed rgp120 in vaccinated guinea pigs. 17b was used as a control to show A32-bound rgp120 had enhanced binding to this CD4-inducible MAb. Liao *et al.* [2004]
- 17b: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of 17b was decreased by trypsin, but increased by thrombin. Thrombin may increase HIV-induced cell fusion in blood by causing a conformational activating shift in gp120. Ling *et al.* [2004]
- 17b: Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The loss of the glycan within the V3 loop (GM299 V3) and two sites adjacent to V3, C2 (GM292 C2) and (GM329 C3), increased neutralization susceptibility to CD4i FAb X5, but each of the glycan mutants and SF162 were refractive to neutralization with 48d and 17b. The loss of sites in C4 (GM438 C4), or V5 (GM454 V5) did not increase neutralization susceptibility to FAb X5. V3 glycans tended to shield V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans shielded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004]
- 17b: A set of HIV-1 chimeras that altered V3 net charge and glycosylation patterns in V1V2 and V3, involving inserting V1V2 loops from a late stage primary isolate taken after the R5 to X4 switch, were studied with regard to phenotype, coreceptor usage, and MAb neutralization. The loops were cloned

into a HXB2 envelope with a LAI viral backbone. It was observed that the addition of the late-stage isolate V1V2 region and the loss of V3-linked glycosylation site in the context of high positive charge gave an X4 phenotype. R5X4, R5, and X4 viruses were generated, and sCD4, 2G12 and b12 neutralization resistance patterns were modified by addition of the late stage V1V2, glycosylation changes, and charge in concert, while neutralization by 2F5 was unaffected. 15e, 17b, and 48d could not neutralize any of the variants tested. Nabatov *et al.* [2004]

- 17b: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including 17b. Pantophlet *et al.* [2004]
- 17b: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12 which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Three CD4i MAbs were tested; all preferentially neutralized SF162, and JRFL became neutralization sensitive to CD4i Abs if the SF162 V1V2 loop was exchanged. Pinter et al. [2004]
- 17b: The SOS mutant envelope protein introduces a covalent disulfide bond between gp120 surface and gp41 transmembrane proteins into the R5 isolate JR-FL by adding cysteines at residues 501 and 605. Pseudovirions bearing this protein bind to CD4 and co-receptor bearing cells, but do not fuse until treatment with a reducing agent, and are arrested prior to fusion after CD4 and co-receptor engagement. CD4i Abs 17b and X5 were weakly neutralizing in all formats, WT, SOS, and when added postbinding. Binley et al. [2003]
- 17b: Called 1.7b. The MAb B4e8 binds to the base of the V3 loop, neutralizes multiple primary isolates and was studied for interaction with other MAbs. B4e8 enhanced binding of CD4i MAbs 4.8d, 1.7b, and A1g8 to R5X4 virus 92HT593, but only of 48d to the R5 virus 92US660, and there was only a modest impact of the combination of B4e8 and CD4i MAbs on neutralization. Cavacini *et al.* [2003]
- 17b: A sCD4-17b single chain chimera was made that can bind to the CD4 binding site, then bind and block co-receptor interaction. This chimeric protein is a very potent neutralizing agent, more potent than IgG1b12, 2G12 or 2F5 against Ba-L infection of CCR5-MAGI cells. It has potential for prophylaxis or therapy. It neutralized 5/6 R5 and X4 strains from the B clade, but was only moderately protective against a D clade isolate, and did not neutralize clade A, C, E, and F isolates. Dey et al. [2003]
- 17b: 17b is known to be comprised of elements from four discontinuous beta strands. Using 17b MAb to select peptides from a combinatorial library, and analyzing the peptides using

- a novel discontinuous epitope reconstruction program, enabled epitope prediction. Segments of gp120 were reconstructed as an antigenic protein mimetic recognized by 17b. Comparisons then were made with a similar prediction of contact residues for CG10, a CD4i MAb that competes with 17b, but has a distinct binding site. Enshell-Seijffers *et al.* [2003]
- 17b: Vaccination of a gp120-CD4 fusion complex in six transgenic XMG2 XenoMouse mice that produce human IgG2 with K light chain did not produce any neutralizing antibodies. 36/39 MAbs derived from one of these mice were in one of two competition groups that were conformational and specific for the complex, suggesting this chimeric vaccine may be of little value, as immunodominant responses recognized epitopes not present in native Env. MAbs from the two CD4-gp120 complex-specific competition groups did not compete with MAbs with known targets on HIV-1 gp120, but their binding was enhanced by binding of 17b. He et al. [2003]
- 17b: This study shows the fragments of CD4i MAbs are better able to neutralize virus than whole IgG. Neutralization of HIV-1 R5 isolates JRFL, JR-CSF and ADA by CD4i MAbs X5, 17b, and 48d decreased with increased molecule size, the neutralizing potency of single-chain Fv (scFv) > than Fab fragments > whole Ab molecules. (With the exception of IgG 48d neutralization of HIV-1 ADA.) HIV-1 X4 isolates 89.6 and HxB2 are both relatively sensitive even to the larger IgG version. R5X4 isolate neutralization was dependent on the isolate and co-receptor usage. The CD4i MAb fragments neutralize HIV-1 subsequent to CD4 binding. The CD4i MAbs bind near the co-receptor binding sites on gp120. Co-receptors bind to the conserved beta19 strand and part of the V3 loop, regions that are masked by the V1V2 loops in the CD4-unbound state. When CD4 is bound, the co-receptor site is exposed near the membrane surface where it would be optimally accessible to co-receptors, and the smaller versions of the molecules are better able to overcome the steric hindrence. Labrijn et al. [2003]
- 17b: Env genes derived from uncultured brain biopsy samples from four HIV-1 infected patients with late-stage AIDS were compared to env genes from PBMC samples. Brain isolates did not differ in the total number or positions of N-glycosylation sites, patterns of coreceptor usage, or ability to be recognized by gp160 and gp41 MAbs. 17b recognized most variants, some from each of the four individuals, by gp120 immunoprecipitation. Ohagen et al. [2003]
- 17b: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional Nlinked glycosylation site sequons inhibiting binding of nonneutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b]
- 17b: This paper describes the generation of CD4i MAb E51, that like CD4i MAb 17b, blocks CCR5 binding to sCD4-bound gp120. E51 has more cross-neutralizing potency than other prototype CD4i MAbs (17b) for B and C clade isolates. E51 and 17b both neutralized HIV-1 clade B strains HXBc2 and ADA, while JR-FL and 89.6 were only neutralized by E51, not 17b. Clade C strains MCGP1.3 and SA32 were both inhibited

by 17b and E51, but E51 was more potent against SA32. The substitutions E381R, F383S, R419D I420R, K421D, Q422L, I423S, and Y435S (HXB2 numbering) all severely reduce 17b and E51 binding. All but I423S also diminish CCR5 binding by more than 50%. The mutation F383S also inhibits sCD4 binding and F105 binding, and K421D inhibits F105 binding, but not sCD4. Xiang *et al.* [2003]

- 17b: The HIV-1 primary isolate DH012 has preserved the epitopes for the MAbs IgG1b12, 2G12, 17b, however natural DH012 infection in chimpanzees and DH012 gp120 vaccination in guinea pigs does not give rise to Abs against these epitopes. Zhu *et al.* [2003]
- 17b: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a CD4i MAb that had no impact on 4KG5 binding. Zwick *et al.* [2003]
- 17b: The two N-terminal domains of CD4, termed D1 and D2, when expressed in the absence of the remaining domains of CD4 retain the capacity to bind to gp120—coding sequences of D1D2 and Igαtp were fused to create a large, multivalent rec protein D1D2Igαtp, which, unlike CD4, does not enhance infection at sub-optimal concentrations—the MAb 17b can also enhance viral replication at sub-optimal concentrations, but D1D2-Igα inhibited the 17b enhancement of two primary isolates. Arthos *et al.* [2002]
- 17b: gp120 mutants were used to define the CXCR4 binding site using CXCR4 displayed on paramagnetic proteoliposomes (PMPLs) to reduce non-specific gp120 binding—basic residues in the V3 loop and the β19 strand (RIKQ, positions 419-422) were involved, and deletion of the V1-V2 loops allowed CD4-independent CXCR4 binding—MAbs 17b (CD4i) and F105 (CD4BS) were used to study conformational changes in the mutants—the affinity of ΔV1 and ΔV1-V2 for 17b was dramatically increased and no longer inducible in the presence of sCD4—V3 mutants R298A and R327A were not recognized by 17b except in the presence of sCD4—mutations in the β19 strand dramatically reduced 17b affinity in the presence or absence of sCD4, consistent with known 17b contact residues in this region. Basmaciogullari et al. [2002]
- 17b: This study examined antibody interactions, binding and neutralization with a B clade R5 isolate (92US660) and R5X4 isolate (92HT593). Abs generally bound and neutralized the R5X4 isolate better than the R5 isolate. Anti-V3 MAb B4a1 increased binding of CD4i MAbs 48d, 17b and A1g8, but only A1g8 binding was increased by B4a1 to the R5 isolate. Additive affects on neutralization of the R5X4 isolate with B4a1 and CD4i MAbs was observed, presumably due to increased exposure of the CD4i binding site, but not for the R5 isolate. Anti-gp41 MAb F240 had a synergistic effect on neutralization with CD4i MAbs 48d and 17b, but not with A1g8 for the R5X4 virus. Cavacini et al. [2002]
- 17b: CD4 residue Phe43 significantly contributes to the affinity of CD4-gp120 interactions – despite decreased affinities for gp120, CD4 proteins and CD4-mimetic peptides lacking a Phe

- side-chain enhance binding of gp120 to 17b in a manner similar to Phe-bearing ligands indicating the Phe42 interaction is not critical for CD4-induced conformational changes in gp120. Dowd *et al.* [2002]
- 17b: Truncation of the gp41 cytoplasmic domain of X4, R5, and X4R5 viruses forces a conformation that more closely resembles the CD4 bound state of the external Envelope, enhancing binding of CD4i MAbs 17b and 48d and of CD4BS MAbs F105, b12, and in most cases of glycosylation site dependent MAb 2G12 and the anti-gp41 MAb 246D in contrast, binding of the anti-V2 MAb 697D and the anti-V3 MAb 694/98D were not affected viruses bearing the truncation were more sensitive to neutralization by MAbs 48d, b12, and 2G12 the anti-C5 MAb 1331A was used to track levels of cell surface expression of the mutated proteins. Edwards *et al.* [2002]
- 17b: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I and Cluster II MAbs required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor, binding to a fusion intermediate. 17b was used to demonstrate that the Cluster I and II MAbs bound to gp120/gp41 complexes, not to gp41 after shedding of gp120. Finnegan *et al.* [2002]
- 17b: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion the preincubation 31.5 C step did not alter the inhibitory activity of neutralizing Abs anti-gp41 2F5, or anti-gp120 2G12, IG1b12, 48d, and 17b. Golding *et al.* [2002b]
- 17b: HIV-1 gp160ΔCT (cytoplasmic tail-deleted) proteoliposomes (PLs) containing native, trimeric envelope glycoproteins from R5 strains YU2 and JRFL, and X4 strain HXBc2, were made in a physiologic membrane setting as candidate immunogens for HIV vaccines—2F5 bound to gp160ΔCT with a reconstituted membrane ten-fold better than the same protein on beads—anti-CD4BS MAbs IgG1b12 and F105, A32 (C1-C4), C11 (C1-C5), and 39F (V3) MAbs bound gp160ΔCT PLs indistinguishably from gp160ΔCT expressed on the cell surface—non-neutralizing MAbs C11 and A32 bound with lower affinity than NAb IgG1b12—the MAb 17b was sCD4 inducible on gp160ΔCT PL. Grundner et al. [2002]
- 17b: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar. Enthalpy and entropy changes were divergent, but compensated. Not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs (17b, 48d, 1.5e, b6, F105 and F91) had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, but the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. The high values suggest surface burial or protein folding an ordering of amino acids. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer),

- a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong *et al.* [2002]
- 17b: A pseudotyping assay showed that an X4 V3 loop peptide could enhance infectivity of X4 virus, R5 and R5X4 V3 loops peptides could enhance infectivity of an R5 virus, and R5X4 peptides could enhance infectivity of an R5X4 virus. Neither R5 nor R5X4 peptides influenced binding of CD4BS MAbs F105 and Ig1Gb12, but did increase binding of CD4i MAb 17b. Ling et al. [2002]
- 17b: Ab binding characteristics of SOS gp140 were tested using SPR and RIPA SOS gp140 is gp120-gp41 bound by a disulfide bond NAbs 2G12, 2F5, IgG1b12, CD4 inducible 17b, and 19b bound to SOS gp140 better than uncleaved gp140 (gp140unc) and gp120 non-neutralizing MAbs 2.2B (binds to gp41 in gp140unc) and 23A (binds gp120) did not bind SOS gp140. Schulke *et al.* [2002]
- 17b: Oligomeric gp140 (o-gp140) derived from R5 primary isolate US4 was characterized for use as a vaccine reagent antigen capture ELISA was used to compare the antigenicity of gp120 and o-gp140 using a panel of well characterized MAbs 17b recognized both gp120 monomer and o-gp140. Srivastava et al. [2002]
- 17b: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations 375 S/W seems to favor a conformation of gp120 closer to the CD4-bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang et al. [2002b]
- 17b: Five CD4i MAbs were studied, 17b, 48d and three new MAbs derived by Epstein-Barr virus transformation of PBMC from an HIV+ long term non-progressor 23e and 21c were converted to hybridomas to increase Ab production all compete with the well-characterized 17b CD4i MAb in an ELISA antigen capture assay critical binding residues are mapped and the CD4i MAb epitopes were distinct but share a common element near isoleucine 420, also important for CCR5 binding, and all five can block CCR5 binding to a sCD4-gp120 complex the MAb 48d has the epitope most similar to the CCR5 binding site. Xiang *et al.* [2002a]
- 17b: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin—stabilized oligomer gp140Δ683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and C11, A32, and 30D which did not bind the stabilized oligomer. Yang et al. [2002]

- 17b: A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang et al. [2002]
- 17b: Using a cell-fusion system, it was found CD4i antibodies 17b, 48d, and CG10 reacted faintly with Env expressing HeLA cells even in the absence of sCD4 or CD4 expressing target cells. Reactivity increased after sCD4 addition, but not after CD4 expressing target cell addition, and binding was not increased at the cell-to-cell CD4-Env interface. This suggests the CD4i co-receptor binding domain is largely blocked at the cell-fusion interface, and so CD4i antibodies would not be able access this site and neutralize cell-mediated viral entry. Finnegan et al. [2001]
- 17b: Mutations in two glycosylation sites in the V2 region of HIV-1 ADA at positions 190 and 197 (187 DNTSYRLINCNTS 199) cause the virus to become CD4-independent and able to enter cells through CCR5 alone—these same mutations tended to increase the neutralization sensitivity of the virus, including to 17b—only the CD4i antibodies 17b and 48d showed an increased affinity of the CD4 independent viruses relative to wild-type. Kolchinsky et al. [2001]
- 17b: Structural aspects of the interaction of neutralizing Abs with HIV-1 Env are reviewed Env essentially has three faces, one is largely inaccessible on the native trimer, and two that exposed but have low immunogenicity on primary viruses neutralization is suggested to occur by inhibition of the interaction between gp120 and the target cell membrane receptors as a result of steric hindrance and it is noted that the attachment of approximately 70 IgG molecules per virion is required for neutralization, which is equivalent to about one IgG molecule per spike the 2G12, 17b and b12 epitopes are discussed in detail the 17b epitope is masked prior to CD4 binding by the V1-V2 loop and in contrast to sCD4, the binding of cell surface CD4 to virus does not appear to make the epitope accessible to binding by 17b to allow neutralization. Poignard *et al.* [2001]
- SHIV-HXBc2 is a neutralization sensitive non-pathogenic virus, and several *in vivo* passages through monkey's yielded highly pathogenic SHIV KU-1 HXBc2 and the KU-1 clone HXBc2P3.2 differ in 12 amino acids in gp160 substitutions in both gp120 and gp41 reduced the ability of sCD4, IgG1b12, F105 and AG1121 to Env achieve saturation and full occupancy, and neutralize KU-1 17b and 2F5 also bound less efficiently to HXBc2P3.2, although 2G12 was able to bind both comparably. Si *et al.* [2001]
- 17b: Abs against the V3 loop (50.1, 58.2, 59.1, 257-D, 268-D, 447-52D), CD4BS (IgG1b12, 559-64D, F105), CD4i (17b), and to gp41 (2F5, F240) each showed similar binding efficiency to Env derived from related pairs of primary and TCLA lines (primary: 168P and 320SI, and TCLA: 168C and 320SI-C3.3), but the TCLA lines were much more susceptible to neutraliza-

tion suggesting that the change in TCLA lines that make them more susceptible to NAbs alters some step after binding – 17b bound at somewhat greater levels to 168C than to 168P, but this is not a general feature of 17b binding to primary versus TCLA strains. York *et al.* [2001]

- 17b: 17b binds to a CD4 inducible epitope which partially overlaps the CCR5 binding site JRFL, YU2, 89.6, and HXB2 and their C1-, V1/V2-, C5 -deletion mutants were used to study how 17b binding affects gp120-CD4 interactions 17b reduced CD4-gp120 interactions by decreasing the on-rate and increasing the off-rate of sCD4, while enhanced binding of sCD4 binding was observed for the 17b-bound, V1/V2 deleted gp120s 17b was considered to be a surrogate for CCR5, and the authors suggest that 17b binding may shift V1/V2 into a position that interferes with CD4 binding, forcing a release. Zhang *et al.* [2001a]
- 17b: To determine the antigenicity of virus killed by thermal and chemical inactivation, retention of conformation-dependent neutralization epitopes was examined, and exposure of CD4BS epitopes was found to be enhanced (MAbs IgG1b12, 205-46-9, and 205-43-1) binding to 2G12 and 447-52D epitopes was essentially unaltered the 17b CD4i epitope was also exposed. Grovit-Ferbas *et al.* [2000]
- 17b: SF162 is a neutralization-resistant HIV-1 isolate N-linked glycosylation modifications in the V2 loop of the SF162 gp120 revealed that these sites prevent neutralization by CD4BS MAbs (IgG1b12 and IgGCD4), and protect against neutralization by V3 MAbs (447-D and 391-95D) V2-region glycosylation site mutations did not alter neutralization resistance to V2 MAbs (G3.4 and G3.136) or CD4i MAbs (17b and 48d) V2 glycosylation site modification allows infection of macrophages, probably due to glycosylated forms requiring fewer CCR5 molecules for viral entry. Ly & Stamatatos [2000]
- 17b: Six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park *et al.* [2000]
- 17b: Mutagenesis defines Ile-420, Lys-421, Gln-422, Pro-438, and Gly-441 to be important residues for CCR5 binding – these positions are located on two strands that connect the gp120 bridging sheet and outer domain, suggesting a mechanism for conformational shifts induced by CD4 binding to facilitate CCR5 binding. Rizzuto & Sodroski [2000]
- 17b: sCD4 can activate fusion between effector cells expressing
   Env and target cells expressing coreceptor (CCR5 or CXCR4)
   alone without CD4 CD4i MAbs 17b and 48d have little
   effect on a standard cell fusion assay but potently block sCD4
   activated fusion 17b was broadly cross-reactive inhibiting
   sCD4 activated fusion with Env from clades A, B, C, D, E, F,
   and F/B. Salzwedel *et al.* [2000]
- 17b: Soluble gp140 derived from SF162, a neutralization-resistant primary isolate, and SF162AV2 a neutralization-susceptible isolate with 30 amino acids deleted from the V2 loop, were generated with or without the gp120-gp41 cleavage site intact all forms are recognized by oligomer-specific MAb T4 and show enhanced binding of CD4i MAb 17b when sCD4 is bound the fused forms are less efficiently recognized than

- the cleaved forms by polyclonal neutralizing sera from HIV-infected patients the V3 loop is more exposed on the fused form. Stamatatos *et al.* [2000]
- 17b: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang *et al.* [2000]
- 17b: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits – a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen - SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 – nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999]
- 17b: A CD4-independent viral variant of IIIB, IIIBx, was generated on CXCR4-expressing cells IIIBx exhibited greater exposure of the 17b and 48d epitopes and enhanced neutralization by CD4i MAbs and by polyclonal human sera the 17b epitope has significant overlap with the CCR5 coreceptor binding site. Hoffman *et al.* [1999]
- 17b: A panel of MAbs was shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a structure closely approximating full length folded monomer CD4i MAbs 17b and 48d bound better to the deleted protein than to wild type. Binley et al. [1998]
- 17b: 17b Fab was co-crystallized with a gp120 core and CD4, and it's binding site can be directly visualized—17b binds to the "bridging sheet" of gp120, an antiparallel beta sheet region, contacting residues from the C4 region and the V1/V2 stem—the contact area is small for an Ab-antigen interactive surface, and dominated in the Ab by the heavy chain—the center of the binding region has hydrophobic interactions, and the periphery charge interactions, acidic on 17b and basic on gp120. Kwong et al. [1998]
- 17b: Moore and Binley provide a commentary on the papers by Rizzuto *et al.* [1998], Wyatt *et al.* [1998] and Kwong *et al.* [1998] they point out 17b shares binding elements in gp120 with chemokine receptor molecules, and that CD4 needs to bind to gp120 first to make the 17b epitope accessible and it

may be sterically blocked in the CD4 bound virus, thus making it a poor NAb for primary isolates Moore & Binley [1998]. Kwong *et al.* [1998]; Moore & Binley [1998]; Rizzuto *et al.* [1998]; Wyatt *et al.* [1998]

- 17b: Site directed mutagenesis of a WU2 protein with the V1-V2 loops deleted revealed key residues for 17b-gp120 interaction and interaction of gp120 and CCR5 mutations in residues that reduced 17b by 70% were R/D 419, I/R 420, Q/L 422, Y/S 435, I/S 423, K/D 121 and K/D 421– 17b can neutralize HIV-1 strains that use different chemokine receptors, supporting a common region in gp120 in chemokine-receptor interaction. Rizzuto et al. [1998]
- 17b: Deleting the V2 loop of neutralization-resistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F deletion of V2 but not V1 enabled neutralization by CD4i MAbs 17b and 48d. Stamatatos & Cheng-Mayer [1998]
- 17b: sCD4 induces 17b binding in primary isolates and TCLA strains amino acids that reduce the efficiency of binding were determined and found also to compromise syncytia formation and viral entry V1V2 deletion or sCD4 binding can expose the 17b epitope for both HXBc2 and macrophage tropic YU2 neutralizing potency of 17b is probably weak due to poor exposure of the epitope 17b epitope exposure upon sCD4 binding can occur over a wide range of temperatures, consistent with the energy of CD4 binding being sufficient to drive the V1/V2 loop into a new conformation. Sullivan et al. [1998b]
- 17b: The HIV-1 virus YU2 entry can be enhanced by MAbs binding to the CD4BS, V3 loop, and CD4i epitopes the activation for this enhanced entry state could be conferred on HxB2 by introducing the YU2 V3 loop, or the YU2 V3 and V1/V2 loops, and the presence of V1/V2 increased the enhancement a similar effect is observed by sub-neutralizing concentrations of sCD4 and the effect is dependent of CCR5 17b enhances YU2 enhanced viral entry 10-fold, whereas HXBc2 was neutralized. Sullivan *et al.* [1998a]
- 17b: Summary of the implications of the crystal structure of a gp120 core bound to CD4 and 17b, combined with what is known about mutations that reduce NAb binding to gp120 - probable mechanism of neutralization is interference with chemokine receptor binding - mutations in 88N, 117K, 121K, 256S, 257T, N262, Delta V3, E370, E381, F 382, R 419, I 420, K 421, Q 422, I 423, W 427, Y 435, P 438, M 475 of HXBc2 (IIIB) reduce binding – the only variable residues in gp120 that contact 17b are 202T and 434M - the contact points for 17b with the crystallized incomplete gp120 are mostly in the heavy chain of the Ab, and there is a gap between 17b's light chain and the partial gp120 which may be occupied by the V3 loop in a complete gp120 molecule – the authors propose that the V2 and V3 loops may mask the CD4i Ab binding site, and that the V2 loop may be repositioned upon CD4 binding. Wyatt et al. [1998]
- 17b: Virus with the V1-V2 loop deleted was viable and more susceptible to neutralization by CD4i MAb 17b, and anti-V3 MAbs 1121, 9284, and 110.4, but not to a CD4BS MAb, F105, or sCD4. Cao *et al.* [1997b]

- 17b: Study shows neutralization is not predicted by MAb binding to JRFL monomeric gp120, but is associated with oligomeric Env binding 17b bound monomer, oligomer, and neutralized JRFL in the presence of sCD4, but if sCD4 was not present, 17b only bound monomer. Fouts *et al.* [1997]
- 17b: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env – 17b has synergistic response in combination with anti-V3 MAb 694/98-D. Li *et al.* [1997]
- 17b: Neutralizes TCLA strains, but not primary isolates. Parren *et al.* [1997b]
- 17b: 48d binds to the IIIB protein and not IIIB V3 peptide, while binding to the Can0A V3 peptide, suggesting Can0A V3 is a conformer that mimics the 48d it does not bind to 17b, distinguishing the epitopes. Weinberg *et al.* [1997]
- 17b: Binds to sgp120 efficiently, but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding partial re-exposure if sCD4 was bound could not bind to HXBc2 gp120 if the 19 C-term amino acids were deleted in conjunction with amino acids 31-93 in C1, but binding was restored in the presence of sCD4. Wyatt *et al.* [1997]
- 17b: Many MAbs inhibit binding (anti-C1, -C5, -C4, -CD4BS)

   anti-V3 MAb 5G11 enhances binding, as do C1-C4 discontinuous epitopes A32 and 2/11c enhances binding of some anti-V2 MAbs. Moore & Sodroski [1996]
- 17b: Binding did not result in significant gp120 dissociation from virion, in contrast to 48d, although the gp41 epitope of MAb 50-69 was exposed. Poignard et al. [1996a]
- 17b: Neutralizes JR-FL inhibits gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola *et al.* [1996a]
- 17b: MIP-1α binding to CCR-5 expressing cells can be inhibited by gp120-sCD4 binding of 17b blocks this inhibition.
   Wu et al. [1996]
- 17b: Binds with higher affinity to monomer and oligomer, slow association rate, poor neutralization of lab strain this is in contrast to 48d, which has very different kinetics. Sattentau & Moore [1995]
- 17b: Studies using a V1/V2 deletion mutant demonstrated that enhanced binding of 17b in the presence sCD4 involves the V1/V2 loops, with more significant involvement of V2 similar effect observed for 48d and A32. Wyatt *et al.* [1995]
- 17b: A mutation in gp41, 582 A/T, confers resistance to neutralization (also confers resistance to MAbs F105, 48d, 21h and 15e). Thali *et al.* [1994]
- 17b: Binding of 48d is much more influenced by sequence variation among molecular clones of LAI than is binding of 17b. Moore *et al.* [1993c]
- 17b: Epitope is better exposed upon CD4 binding to gp120 competes with 15e and 21h, anti-CD4 binding site MAbs 113 D/R, 252 R/W, 257 T/A or G, 370 E/D, 382 F/L, 420 I/R, 433A/L, 438 P/R and 475 M/S confer decreased sensitivity to neutralization. Thali et al. [1993]

No. 1120 MAb ID 21c (2.1C) HXB2 Location Env Author Location gp120 (IIIB, J62) Epitope Subtype B

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

Ab Type gp120 CD4i, gp120 CCR5BS

Research Contact James Robinson, Tulane University, New Or-

leans, LA, USA

References Haynes et al. 2005; Gorny & Zolla-Pazner 2004; Xiang et al. 2002b; Xiang et al. 2002a

**Keywords** antibody binding site definition and exposure, antibody generation, review, vaccine antigen

design

- 21c: Called 2.1C. Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 2.1C has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)
- 21c: This review summarizes MAbs directed to HIV-1 Env. There are six CD4 inducible MAbs and Fabs in the database. The MAb forms neutralize TCLA strains only, but the smaller Fabs and scFv fragments can neutralize primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 21c: Five CD4i MAbs were studied, 17b, 48d and three new MAbs derived by Epstein-Barr virus transformation of PBMC from an HIV+ long term non-progressor – 23e and 21c were converted to hybridomas to increase Ab production - all compete with the well-characterized 17b CD4i MAb in an ELISA antigen capture assay - critical binding residues are mapped and the CD4i MAb epitopes were distinct but share a common element near isoleucine 420, also important for CCR5 binding, and all five can block CCR5 binding to a sCD4-gp120 complex - the MAb 48d has the epitope most similar to the CCR5 binding site. Xiang et al. [2002a] (antibody binding site definition and exposure, antibody generation)
- 21c: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations – 375 S/W seems to favor a conformation of gp120 closer to the CD4bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced -IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced – 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang et al. [2002b] (antibody binding site definition and exposure, vaccine antigen design)

No. 1121

MAb ID 23e

**HXB2 Location** Env

Author Location gp120 (IIIB, J62)

**Epitope** 

Subtype B

Immunogen HIV-1 infection Species (Isotype) human (IgG)

Neutralizing L

Ab Type gp120 CD4i

Research Contact James Robinson, Tulane University, New Or-

leans, LA, USA

References Gorny & Zolla-Pazner 2004; Xiang et al.

2002b; Xiang et al. 2002a

**Keywords** antibody binding site definition and exposure,

antibody generation, review, vaccine antigen

- 23e: This review summarizes MAbs directed to HIV-1 Env. There are six CD4 inducible MAbs and Fabs in the database. The MAb forms neutralize TCLA strains only, but the smaller Fabs and scFv fragments can neutralize primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 23e: Five CD4i MAbs were studied, 17b, 48d and three new MAbs derived by Epstein-Barr virus transformation of PBMC from an HIV+ long term non-progressor - 23e and 21c were converted to hybridomas to increase Ab production - all compete with the well-characterized 17b CD4i MAb in an ELISA antigen capture assay - critical binding residues are mapped and the CD4i MAb epitopes were distinct but share a common element near isoleucine 420, also important for CCR5 binding, and all five can block CCR5 binding to a sCD4-gp120 complex - the MAb 48d has the epitope most similar to the CCR5 binding site. Xiang et al. [2002a] (antibody binding site definition and exposure, antibody generation)
- 23e: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations - 375 S/W seems to favor a conformation of gp120 closer to the CD4bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced -IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced – 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang et al. [2002b] (antibody binding site definition and exposure, vaccine antigen design)

**No.** 1122

**MAb ID** 48d (4.8d, 4.8D)

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

**Neutralizing** L P (weak)

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

**Ab Type** gp120 CD4i

Research Contact James Robinson, Tulane University, New Or-

leans, LA, USA

References Selvarajah et al. 2005; Martín-García et al.

2005; Lusso et al. 2005; Pinter et al. 2004; Pantophlet et al. 2004; Nabatov et al. 2004; McCaffrey et al. 2004; Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2003; Labrijn et al. 2003; Cavacini et al. 2003; Cavacini et al. 2002; Zhang et al. 2002; Edwards et al. 2002; Xiang et al. 2002a; Xiang et al. 2002b; Yang et al. 2002; Golding

et al. 2002b; Kwong et al. 2002; Finnegan et al. 2001; Verrier et al. 2001; Kolchinsky et al. 2001; Salzwedel et al. 2000; Yang et al. 2000; Park et al. 2000; Ly & Stamatatos 2000; Fortin et al. 2000; Hoffman et al. 1999; Oscherwitz et al. 1999a; Stamatatos & Cheng-Mayer 1998; Binley et al. 1998; Yang et al. 1998; Sullivan et al. 1998b; Parren et al. 1998a; Mondor et al. 1998; Wyatt et al. 1998; Frankel et al. 1998; Parren et al. 1997b; Wyatt et al. 1997; Ugolini et al. 1997; Lee et al. 1997; Weinberg et al. 1997; Li et al. 1997; Binley et al. 1997a; Trkola et al. 1996a; Poignard et al. 1996a; Moore & Sodroski 1996; Sattentau & Moore 1995; Sattentau et al. 1995; Wyatt et al. 1995; Sattentau 1995; D'Souza et al. 1995; Moore et al. 1994b; Thali et al. 1994; Moore et al. 1993c; Moore & Ho 1993; Thali et al. 1993

**Keywords** antibody binding site definition and exposure, antibody interactions, binding affinity, coreceptor, kinetics, review, structure, subtype

receptor, kinetics, review, structure, subtype comparisons, vaccine antigen design, vaccinespecific epitope characteristics, variant crossrecognition or cross-neutralization

- 48d: 48d and 17b have similar epitopes, and the pair are unique among human and rodent MAbs.
- 48d: NIH AIDS Research and Reference Reagent Program: 1756.
- 48d: The epitope for the MAb D19 is conserved and embedded in V3. D19 is unique in that for R5 viruses, it was cryptic and did not bind without exposure to sCD4, and for X4 and R5X4 isolates it was constitutively exposed. D19b is unquiue among CD4i antibodies in that it binds to the V3 loop. CD4i MAbs 17b and 48d were used as controls for CD4i characterization; in contrast to D19, other CD4i MAbs bind to the conserved bridging sheet and do not differentiate between R5 and X4 using strains. Lusso *et al.* [2005]
- 48d: The HIV-1 Bori-15 variant was adapted from the Bori isolate for replication microglial cells. Bori-15 had increased replication in microglial cells and a robust syncytium-forming phenotype, ability to use low levels of CD4 for infection, and increased sensitivity to neutralization by sCD4 and 17b. Four amino acid changes in gp120 V1-V2 were responsible for this change. Protein functionality and integrity of soluble, monomeric gp120-molecules derived from parental HIV-1 Bori and microglia-adapted HIV-1 Bori-15 was assessed in ELISA binding assays using F105, IgG1b12, 17b and 48d, 2G12 and 447-52D. Association rates of sCD4 and 17b were not changed, but dissociation rates were 3-fold slower for sCD4 and 14-fold slower for 17b. Equilibrium binding studies showed 48d bound better to Bori-15 than Bori in the absence of sCD4, while 17b bound identically. Martín-García et al. [2005] (antibody binding site definition and exposure)
- 48d: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did

not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened while not obscuring b12 binding. CD4i MAbs (48d, 17b) did not bind to either GDMR or mCHO even with sCD4. Selvarajah *et al.* [2005] (vaccine antigen design, vaccine-specific epitope characteristics)

- 48d: This review summarizes MAbs directed to HIV-1 Env.
  There are six CD4 inducible MAbs and Fabs in the database.
  The MAb forms neutralize TCLA strains only, but the smaller
  Fabs and scFv fragments can neutralize primary isolates. Gorny
  & Zolla-Pazner [2004] (review)
- 48d: Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The loss of the glycan within the V3 loop (GM299 V3) and two sites adjacent to V3, C2 (GM292 C2) and (GM329 C3), increased neutralization susceptibility to CD4i FAb X5, but each of the glycan mutants and SF162 were refractive to neutralization with 48d and 17b. The loss of sites in C4 (GM438 C4), or V5 (GM454 V5) did not increase neutralization susceptibility to FAb X5. V3 glycans tended to shield V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans shielded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004] (antibody binding site definition and exposure, vaccine antigen design)
- · 48d: A set of HIV-1 chimeras that altered V3 net charge and glycosylation patterns in V1V2 and V3, involving inserting V1V2 loops from a late stage primary isolate taken after the R5 to X4 switch, were studied with regard to phenotype, coreceptor usage, and MAb neutralization. The loops were cloned into a HXB2 envelope with a LAI viral backbone. It was observed that the addition of the late-stage isolate V1V2 region and the loss of V3-linked glycosylation site in the context of high positive charge gave an X4 phenotype. R5X4, R5, and X4 viruses were generated, and sCD4, 2G12 and b12 neutralization resistance patterns were modified by addition of the late stage V1V2, glycosylation changes, and charge in concert, while neutralization by 2F5 was unaffected. 15e, 17b, and 48d could not neutralize any of the variants tested. Nabatov et al. [2004] (antibody binding site definition and exposure, co-receptor)
- 48d: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including 48d. Pantophlet et al. [2004] (vaccine antigen design)
- 48d: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i,

(except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Three CD4i MAbs were tested; all preferentially neutralized SF162, and JRFL became neutralization sensitive to CD4i Abs if the SF162 V1V2 loop was exchanged. Pinter *et al.* [2004] (variant cross-recognition or cross-neutralization)

- 48d: Called 4.8d. The MAb B4e8 binds to the base of the V3 loop, neutralizes multiple primary isolates and was studied for interaction with other MAbs. B4e8 enhanced binding of CD4i MAbs 4.8d, 1.7b, and A1g8 to R5X4 virus 92HT593, but only of 48d to the R5 virus 92US660, and there was only a modest impact of the combination of B4e8 and CD4i MAbs on neutralization. Cavacini *et al.* [2003] (antibody interactions, co-receptor)
- 48d: This study shows the fragments of CD4i MAbs are better able to neutralize virus than whole IgG. Neutralization of HIV-1 R5 isolates JRFL, JR-CSF and ADA by CD4i MAbs X5, 17b, and 48d decreased with increased molecule size, the neutralizing potency of single-chain Fv (scFv) > than Fab fragments > whole Ab molecules. (With the exception of IgG 48d neutralization of HIV-1 ADA being better than the Fab – for 48d, only the IgG and Fab forms were available, not the scFv.) HIV-1 X4 isolates 89.6 and HxB2 are both relatively sensitive even to the larger IgG version. R5X4 isolate neutralization was dependent on the isolate and co-receptor usage. The CD4i MAb fragments neutralize HIV-1 subsequent to CD4 binding. The CD4i MAbs bind near the co-receptor binding sites on gp120. Co-receptors bind to the conserved beta19 strand and part of the V3 loop, regions that are masked by the V1V2 loops in the CD4-unbound state. When CD4 is bound, the co-receptor site is exposed near the membrane surface where it would be optimally accessible to co-receptors, and the smaller versions of the molecules are better able to overcome the steric hindrence. Labrijn et al. [2003] (antibody binding site definition and exposure, co-receptor, variant cross-recognition or cross-neutralization)
- 48d: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of non-neutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet *et al.* [2003b] (vaccine antigen design)
- 48d: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a CD4i MAb that had no impact on 4KG5 binding. Zwick et al. [2003] (antibody interactions)
- 48d: This study examined antibody interactions, binding and neutralization with a B clade R5 isolate (92US660) and R5X4 isolate (92HT593). Abs generally bound and neutralized the

R5X4 isolate better than the R5 isolate. Anti-V3 MAb B4a1 increased binding of CD4i MAbs 48d, 17b and A1g8, but only A1g8 binding was increased by B4a1 to the R5 isolate. Additive affects on neutralization of the R5X4 isolate with B4a1 and CD4i MAbs was observed, presumably due to increased exposure of the CD4i binding site, but not for the R5 isolate. Anti-gp41 MAb F240 had a synergistic effect on neutralization with CD4i MAbs 48d and 17b, but not with A1g8 for the R5X4 virus. Cavacini *et al.* [2002] (variant cross-recognition or cross-neutralization)

- 48d: Truncation of the gp41 cytoplasmic domain of X4, R5, and X4R5 viruses forces a conformation that more closely resembles the CD4 bound state of the external Envelope, enhancing binding of CD4i MAbs 17b and 48d and of CD4BS MAbs F105, b12, and in most cases of glycosylation site dependent MAb 2G12 and the anti-gp41 MAb 246D in contrast, binding of the anti-V2 MAb 697D and the anti-V3 MAb 694/98D were not affected viruses bearing the truncation were more sensitive to neutralization by MAbs 48d, b12, and 2G12 the anti-C5 MAb 1331A was used to track levels of cell surface expression of the mutated proteins. Edwards *et al.* [2002] (**co-receptor**)
- 48d: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion the preincubation 31.5 C step did not alter the inhibitory activity of neutralizing Abs anti-gp41 2F5, or anti-gp120 2G12, IG1b12, 48d, and 17b. Golding *et al.* [2002b]
- 48d: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar. Enthalpy and entropy changes were divergent, but compensated. Not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs (17b, 48d, 1.5e, b6, F105 and F91) had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, but the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. The high values suggest surface burial or protein folding an ordering of amino acids. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong et al. [2002] (antibody binding site definition and exposure)
- 48d: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations 375 S/W seems to favor a conformation of gp120 closer to the CD4-bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced IgG1b12 failed to neutralize this mutant, while neutralization

by 2G12 was enhanced – 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope – another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang *et al.* [2002b]

- 48d: Five CD4i MAbs were studied, 17b, 48d and three new MAbs derived by Epstein-Barr virus transformation of PBMC from an HIV+ long term non-progressor 23e and 21c were converted to hybridomas to increase Ab production all compete with the well-characterized 17b CD4i MAb in an ELISA antigen capture assay critical binding residues are mapped and the CD4i MAb epitopes were distinct but share a common element near isoleucine 420, also important for CCR5 binding, and all five can block CCR5 binding to a sCD4-gp120 complex the MAb 48d has the epitope most similar to the CCR5 binding site. Xiang et al. [2002a] (antibody binding site definition and exposure, co-receptor)
- 48d: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin – stabilized oligomer gp140 delta683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and C11, A32, and 30D which did not bind the stabilized oligomer. Yang *et al.* [2002]
- 48d: Called 4.8D A rare mutation in the neutralization sensitive R2-strain in the proximal limb of the V3 region caused Env to become sensitive to neutralization by MAbs directed against the CD4 binding site (CD4BS), CD4-induced (CD4i) epitopes, soluble CD4 (sCD4), and HNS2, a broadly neutralizing sera 2/12 anti-V3 MAbs tested (19b and 694/98-D) neutralized R2, as did 2/3 anti-CD4BS MAbs (15e and IgG1b12), 2/2 CD4i MAbs (17b and 4.8D), and 2G12 and 2F5 thus multiple epitopes on R2 are functional targets for neutralization and the neutralization sensitivity profile of R2 is intermediate between the highly sensitive MN-TCLA strain and the typically resistant MN-primary strain. Zhang *et al.* [2002] (variant cross-recognition or cross-neutralization)
- 48d: Using a cell-fusion system, it was found CD4i antibodies 17b, 48d, and CG10 reacted faintly with Env expressing HeLA cells even in the absence of sCD4 or CD4 expressing target cells. Reactivity increased after sCD4 addition, but not after CD4 expressing target cell addition, and binding was not increased at the cell-to-cell CD4-Env interface. This suggests the CD4i co-receptor binding domain is largely blocked at the cell-fusion interface, and so CD4i antibodies would not be able access this site and neutralize cell-mediated viral entry. Finnegan *et al.* [2001] (antibody binding site definition and exposure)
- 48d: Mutations in two glcosylation sites in the V2 region of HIV-1 ADA at positions 190 and 197 (187 DNTSYRLINCNTS 199) cause the virus to become CD4-independent and able to enter cells through CCR5 alone – these same mutations tended to increase the neutralization sensitivity of the virus, including to 48d – only the CD4i antibodies 17b and 48d showed an

- increased affinity of the CD4 independent viruses relative to wild-type. Kolchinsky *et al.* [2001]
- 48d: Called 4.8d A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001]
- 48d: Called 4.8D host encoded intercellular adhesion molecule (ICAM-1) is incorporated by the HIV-1 virion and enhances viral infectivity ICAM-1 does not modify virus sensitivity to antibodies 0.5beta or 4.8D or sCD4, but neutralizing ability of F105 was diminished in ICAM bearing virions in the presence of lymphocyte function-association antigen-1 (LFA-1) Ab. Fortin et al. [2000]
- 48d: SF162 is a neutralization-resistant HIV-1 isolate N-linked glycosylation modifications in the V2 loop of the SF162 gp120 revealed that these sites prevent neutralization by CD4BS MAbs (IgG1b12 and IgGCD4), and protect against neutralization by V3 MAbs (447-D and 391-95D) V2-region glycosylation site mutations did not alter neutralization resistance to V2 MAbs (G3.4 and G3.136) or CD4i MAbs (17b and 48d) V2 glycosylation site modification allows infection of macrophages, probably due to glycosylated forms requiring fewer CCR5 molecules for viral entry. Ly & Stamatatos [2000]
- 48d: Six mutations in MN change the virus from a high-infectivity neutralization resistant phenotype to low-infectivity neutralization sensitive V3, CD4BS, and CD4i MAbs are 20-100 fold more efficient at neutralizing the sensitive form the mutation L544P reduced binding of all MAbs against gp120 by causing conformational changes. Park et al. [2000]
- 48d: sCD4 can activate fusion between effector cells expressing Env and target cells expressing coreceptor (CCR5 or CXCR4) alone without CD4 – CD4i MAbs 17b and 48d have little effect on a standard cell fusion assay but potently block sCD4 activated fusion. Salzwedel *et al.* [2000] (co-receptor)
- 48d: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) that preserve and expose some neutralizing epitopes while occluding some non-neutralizing epitopes CD4BS MAbs (F105 and F91) and CD4i (17b and 48d) recognized gp140-GNC4 as well as gp120 or gp140 non-neutralizing MAbs C11, A32, 522-149, M90, and #45 bound to the gp140-GNC4 glycoprotein at reduced levels compared to gp120 MAbs directed at the extreme termini of gp120 C1 (135/9 and 133/290) and C5 (CRA-1 and M91) bound efficiently to gp140-GNC4. Yang *et al.* [2000]
- 48d: A CD4-independent viral variant of IIIB, IIIBx, was generated on CXCR4-expressing cells IIIBx exhibited greater exposure of the 17b and 48d epitopes and enhanced neutralization by CD4i MAbs and by polyclonal human sera. Hoffman et al. [1999]
- 48d: A panel of MAbs were shown to bind with similar or greater affinity and similar competition profiles to a deglycosylated or variable loop deleted core gp120 protein ( Delta V1, V2, and V3), thus such a core protein produces a struc-

ture closely approximating full length folded monomer – CD4i MAbs 17b and 48d bound better to the deleted protein than to wild type. Binley *et al.* [1998]

- 48d: Infection of dendritic cells cultured from CD14+ blood cells or from cadaveric human skin was blocked by neutralizing MAbs IgG1b12, or 2F5 and 2G12 delivered together, but not by control non-neutralizing anti-gp120 MAb 4.8D, indicating that NAbs could interrupt early mucosal transmission events. Frankel et al. [1998]
- 48d: Inhibits binding of Hx10 to both CD4 positive and CD4 negative HeLa cells. Mondor *et al.* [1998]
- 48d: The MAb and Fab binding to the oligomeric form of gp120 and neutralization were highly correlated – authors suggest that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren *et al.* [1998a]
- 48d: Deleting the V2 loop of neutralization-resistant HIV-1 isolate SF162 does not abrogate its replication in PBMC or macrophages, but it enhances its neutralization sensitivity to sera from patients with B clade infection up to 170-fold, and also enhances sensitivity to sera from clades A through F deletion of V2 but not V1 enabled neutralization by CD4i MAbs 17b and 48d. Stamatatos & Cheng-Mayer [1998]
- 48d: CD4i MAbs 17b and 48d compete with MAb CG10, and the binding sites may overlap – MAb A32 enhances binding of 17b, 48d and CG10. Sullivan *et al.* [1998b]
- 48d: Summary of the implications of the crystal structure of the core of gp120 bound to CD4 and 17b with what is known about mutations that reduce NAb binding probable mechanism of neutralization of 48d is interference with chemokine receptor binding CD4 binding increases exposure of epitope due to V2 loop movement 88N, 117K, 121K, 256S, 257T, N262, delta V3, E370, E381, F 382, R 419, I 420, K 421, Q 422, I 423, W 427, Y 435, P 438, M 475 mutations in HXBc2 (IIIB) decrease binding. Wyatt et al. [1998] (structure)
- 48d: A neutralization assay was developed based on heminested PCR amplification of the LTR (HNPCR) LTR-HNPCR consistently revealed HIV DNA and was shown to be a rapid, specific and reliable neutralization assay based on tests with 6 MAbs and 5 isolates. Yang et al. [1998]
- 48d: Prefers CD4-gp120 complex to gp120 alone, but does not enhance fusion, in contrast to MAb CG10, in fact it inhibits syncytium formation. Lee *et al.* [1997] (antibody binding site definition and exposure)
- 48d: One of 14 human MAbs tested for ability to neutralize a chimeric SHIV-vpu+, which expressed HIV-1 IIIB env all Ab combinations tested showed synergistic neutralization 48d has synergistic response with MAbs 694/98-D (anti-V3) and F105. Li *et al.* [1997] (antibody interactions)
- 48d: Neutralizes TCLA strains, but not primary isolates. Parren et al. [1997b] (variant cross-recognition or cross-neutralization)
- 48d: Viral binding inhibition by 48d was strongly correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5). Ugolini et al. [1997]
- 48d: 48d binds to the IIIB protein and not IIIB V3 peptide, while binding to the Can0A V3 peptide, suggesting Can0A V3 is a conformer that mimics the 48d, (but not 17b), epitope.

Weinberg *et al.* [1997] (antibody binding site definition and exposure)

- 48d: Binds efficiently to sgp120 but not soluble gp120+gp41, suggesting its gp120 epitope is blocked by gp41 binding. Wyatt *et al.* [1997] (antibody binding site definition and exposure)
- 48d: Many MAbs inhibit binding (anti-C1, -C5, -C4, -CD4BS)

   anti-C1-C4 discontinuous epitope MAbs A32 and 2/11c enhance binding reciprocal enhanced binding with some anti-V2 MAbs. Moore & Sodroski [1996] (antibody interactions)
- 48d: Binding resulted in gp120 dissociation from virion, mimicking sCD4, and exposure of the gp41 epitope of MAb 50-69, in contrast to CD4BS MAbs. Poignard et al. [1996a] (antibody interactions)
- 48d: Neutralizes JR-FL slightly inhibits gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study. Trkola et al. [1996a] (antibody binding site definition and exposure, co-receptor)
- 48d: Called 4.8D Found to neutralize MN, but not JRCSF, two B subtype primary isolates, or a D subtype primary isolate, by most labs in a multi-laboratory study involving 11 labs.
   D'Souza et al. [1995] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 48d: Formalin inactivation of virus at 0.1% formalin for 10 hours at 4 degrees was optimal for inactivation of virus while maintaining epitope integrity. Sattentau *et al.* [1995] (vaccine antigen design)
- 48d: Binds with similar affinity to monomer and oligomer, moderate association rate, potent neutralization – this is in contrast to 17b, which has very different kinetics. Sattentau & Moore [1995] (antibody binding site definition and exposure, kinetics, binding affinity)
- 48d: Studies using a V1/V2 deletion mutant demonstrated that enhanced binding of 48d in the presence of sCD4 involves the V1/V2 loops, with more significant involvement of V2 similar effect observed for 17b and A32. Wyatt *et al.* [1995] (vaccine antigen design)
- 48d: Poor cross-reactivity with gp120 from most clades. Moore et al. [1994b] (subtype comparisons)
- 48d: A mutation in gp41, 582 A/T, confers resistance to neutralization (also confers resistance to MAbs F105, 21h, 15e and 17b). Thali *et al.* [1994] (variant cross-recognition or cross-neutralization)
- 48d: Called 4.8d Neutralizes IIIB reactive with SF-2 gp120 does not inhibit HIV-1 sera from binding to IIIB gp120. Moore & Ho [1993] (variant cross-recognition or cross-neutralization)
- 48d: Binding of 48d is much more influenced by sequence variation among molecular clones of LAI than is binding of 17b. Moore et al. [1993c] (variant cross-recognition or crossneutralization)
- 48d: Epitope is better exposed upon CD4 binding to gp120 competes with ICR 39.13, 15e and 21h, anti-CD4 binding site MAbs inhibited by anti-CD4BS MAb ICR 39.13g and linear anti-C4 MAbs G3-42 and G3-508 113 D/R, 252 R/W, 257 T/A or G, 370 E/D, 382 F/L, 420 I/R, 421 K/L, 433A/L, 438 P/R and 475 M/S confer decreased sensitivity to neutralization. Thali *et al.* [1993] (antibody binding site definition and exposure, antibody interactions)

**No.** 1123

MAb ID 49e

**HXB2 Location** Env

Author Location gp120 (IIIB, J62)

**Epitope** Subtype B

Neutralizing L

Immunogen HIV-1 infection Species (Isotype) human (IgG)

Ab Type gp120 CD4i

Research Contact James Robinson, Tulane University, New Or-

leans, LA, USA

References Gorny & Zolla-Pazner 2004; Xiang et al.

2002b; Xiang et al. 2002a

**Keywords** antibody binding site definition and exposure, antibody generation, review, vaccine antigen

design

- 49e: This review summarizes MAbs directed to HIV-1 Env. There are six CD4 inducible MAbs and Fabs in the database. The MAb forms neutralize TCLA strains only, but the smaller Fabs and scFv fragments can neutralize primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 49e: Five CD4i MAbs were studied, 17b, 48d and three new MAbs derived by Epstein-Barr virus transformation of PBMC from an HIV+ long term non-progressor – 23e and 21c were converted to hybridomas to increase Ab production - all compete with the well-characterized 17b CD4i MAb in an ELISA antigen capture assay - critical binding residues are mapped and the CD4i MAb epitopes were distinct but share a common element near isoleucine 420, also important for CCR5 binding, and all five can block CCR5 binding to a sCD4-gp120 complex - the MAb 48d has the epitope most similar to the CCR5 binding site. Xiang et al. [2002a] (antibody binding site definition and exposure, antibody generation)
- 49e: A series of mutational changes were introduced into the YU2 gp120 that favored different conformations - 375 S/W seems to favor a conformation of gp120 closer to the CD4bound state, and is readily bound by sCD4 and CD4i MAbs (17b, 48d, 49e, 21c and 23e) but binding of anti-CD4BS MAbs (F105, 15e, IgG1b12, 21h and F91 was markedly reduced -IgG1b12 failed to neutralize this mutant, while neutralization by 2G12 was enhanced - 2F5 did not neutralize either WT or mutant, probably due to polymorphism in the YU2 epitope another mutant, 423 I/P, disrupted the gp120 bridging sheet, favored a different conformation and did not bind CD4, CCR5, or CD4i antibodies, but did bind to CD4BS MAbs. Xiang et al. [2002b] (antibody binding site definition and exposure, vaccine antigen design)

No. 1124

MAb ID Fbb21

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4i References Zwick et al. 2003 **Keywords** antibody interactions • Fbb21: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This is a novel CD4i Fab first used in this study. Fbb21, like other CD4i MAbs, did not inhibit or enhance 4KG5 binding. Zwick et al. [2003] (antibody interactions)

No. 1125

MAb ID Fbb21

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4i

References Zwick et al. 2003

Keywords antibody interactions

• Fbb21: scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This is a novel CD4i Fab first used in this study. Fbb21, like other CD4i MAbs, did not inhibit or enhance 4KG5 binding. Zwick et al. [2003] (antibody interactions)

No. 1126

MAb ID X5 (Fab X5)

HXB2 Location Env

Author Location gp120 (JRFL)

**Epitope** 

Subtype B

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 CD4i

References Pinter et al. 2004; Pantophlet et al. 2004; Mc-Caffrey et al. 2004; Darbha et al. 2004; Binley et al. 2004; Gorny & Zolla-Pazner 2004; Pantophlet et al. 2003b; Zwick et al. 2004; Zwick et al. 2003; Zhang et al. 2003; Labrijn et al. 2003; Binley et al. 2003; Moulard et al. 2002

Keywords antibody binding site definition and exposure, antibody generation, antibody interactions, coreceptor, review, structure, subtype comparisons, vaccine antigen design, variant crossrecognition or cross-neutralization

- X5: 93 viruses from different clades were tested for their neutralization cross-reactivity using a panel of HIV antibodies. X5 is a CD4i antibody and neutralized only the most sensitive B-clade envelopes in the pseudovirus assay, but was able to neutralize 2/25 non-B isolates in the PBMC assay, possibly due to differential corecptor expression. Binley et al. [2004] (variant cross-recognition or cross-neutralization, subtype comparisons)
- X5: The structure of the Fab X5 was determined at 1.9 angstrom resolution. The binding site is a long, 22 amino acid CDR H3 with a hook shape. Long CDR H3s are also found in IgG1b12 (18 residues) and 17b (19 residues). FAb X5 has a W100, F100Y in the CDR H3 hook shown to be important for binding through site specific mutagenesis. Compared to JRCSF, Ala substitutions at eight residues reduced binding more than 3 fold: C119, K207, G367, M426, W427, V430, I423, and K432. Only I423A and K432A were thought to possibly directly interact with X5, the other mutations were thought likely to disrupt the overall structure or CD4 binding. Darbha *et al.* [2004] (antibody binding site definition and exposure, structure)
- X5: This review summarizes MAbs directed to HIV-1 Env.
  There are six CD4 inducible MAbs and Fabs in the database.
  The MAb forms neutralize TCLA strains only, but the smaller
  Fabs and scFv fragments can neutralize primary isolates. Gorny
  & Zolla-Pazner [2004] (review)
- X5: Sera from two HIV+ people and a panel of MAbs were used to explore susceptibility to neutralization in the presence or absence of glycans within or adjacent to the V3 loop and within the C2, C4 and V5 regions of HIV-1 SF162 env gp120. The loss of the glycan within the V3 loop (GM299 V3) and two sites adjacent to V3, C2 (GM292 C2) and (GM329 C3), increased neutralization susceptibility to CD4i FAb X5, but each of the glycan mutants and SF162 were refractive to neutralization with 48d and 17b. The loss of sites in C4 (GM438 C4), or V5 (GM454 V5) did not increase neutralization susceptibility to FAb X5. V3 glycans tended to shield V3 loop, CD4 and co-receptor MAb binding sites, while C4 and V5 glycans shielded V3 loop, CD4, gp41 but not co-receptor MAb binding sites. Selective removal of glycans from a vaccine candidate may enable greater access to neutralization susceptible epitopes. McCaffrey et al. [2004] (antibody binding site definition and exposure, vaccine antigen design)
- X5: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including X5. Pantophlet et al. [2004] (vaccine antigen design)
- X5: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and

- 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Three CD4i MAbs were tested; all preferentially neutralized SF162, and JRFL became neutralization sensitive to CD4i Abs if the SF162 V1V2 loop was exchanged. FAb X5 could neutralize both viruses, but had reduced potency against JRFL. Pinter *et al.* [2004] (variant cross-recognition or cross-neutralization)
- X5: Called Fab X5. This paper is a study of the 2F5 NAb complexed to peptide ELDKWAS; the peptide was found to interact with amino acids near the base of the very long (22 residue) CDR 3H region of the Ab, although a Phe at the apex of the loop was also important. The authors suggest that particularly long CDR H3 regions may be a common feature of HIV-1 neuturalizing antibodies there are 22 residues in 2F5's H3, 18 in b12's H3, and 22 residues in X5's H3. They express concern that because small animals like mice are unable to elicit Ab responses with such long H3s, they may be poor model systems for HIV vaccine studies. Zwick *et al.* [2004] (antibody interactions)
- X5: The SOS mutant envelope protein introduces a covalent disulfide bond between gp120 surface and gp41 transmembrane proteins into the R5 isolate JR-FL by adding cysteines at residues 501 and 605. Pseudovirions bearing this protein bind to CD4 and co-receptor bearing cells, but do not fuse until treatment with a reducing agent, and are arrested prior to fusion after CD4 and co-receptor engagement. CD4i Abs X5 and 17b were weakly neutralizing in all formats, WT, SOS, and when added postbinding. Binley *et al.* [2003] (vaccine antigen design)
- X5: This study shows the fragments of CD4i MAbs are better able to neutralize virus than whole IgG. Neutralization of HIV-1 R5 isolates JRFL, JR-CSF and ADA by CD4i MAbs X5, 17b, and 48d decreased with increased molecule size, the neutralizing potency of single-chain Fv (scFv) > than Fab fragments > whole Ab molecules. (With the exception of IgG 48d neutralization of HIV-1 ADA.) HIV-1 X4 isolates 89.6 and HxB2 are both relatively sensitive even to the larger IgG version. R5X4 isolate neutralization was dependent on the isolate and co-receptor usage. The CD4i MAb fragments neutralize HIV-1 subsequent to CD4 binding. The CD4i MAbs bind near the co-receptor binding sites on gp120. Co-receptors bind to the conserved beta19 strand and part of the V3 loop, regions that are masked by the V1V2 loops in the CD4-unbound state. When CD4 is bound, the co-receptor site is exposed near the membrane surface where it would be optimally accessible to co-receptors, and the smaller versions of the molecules are better able to overcome the steric hindrence. Labrijn et al. [2003] (antibody binding site definition and exposure, co-receptor, variant cross-recognition or cross-neutralization)
- X5: A gp120 molecule was designed to focus the immune response onto the IgG1b12 epitope. Ala substitutions that enhance the binding of IgG1b12 and reduce the binding of non-neutralizing MAbs were combined with additional N-linked glycosylation site sequons inhibiting binding of non-neutralizing MAbs; b12 bound to the mutated gp120. C1 and C5 were also removed, but this compromised b12 binding. Pantophlet et al. [2003b] (vaccine antigen design)
- X5: The Fab m18 was selected from a human phage display

library by a new method called sequential antigen panning (SAP), using a series of antigens to screen the library to pick broadly cross-reactive isolates. The ability to block cell mediated fusion by m18 was compared to Fabs X5 and b12 for a clade A, CRF01 EA, G, and 6 clade B isolates, and the inhibitory activity of m18 was slightly lower but comparable to neutralizing Fabs b12 and X5. It also showed broad crossneutralization; 11/15 pseudotyped Envs from primary isolates from clades A-F were inhibited in an IC50 assay at concentration less than or equal to 100 ug/ml; X5 was also tested and somewhat more potent, generally requiring lower concentrations and inhibiting 13/15 primary isolates. Zhang *et al.* [2003] (variant cross-recognition or cross-neutralization, subtype comparisons)

- X5: scFv 4KG5 reacts with a conformational epitope. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120. MAbs to the V2 loop, V3 loop, V3-C4 region, and CD4BS diminished binding, while MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected. This is a CD4i MAb that had no impact on 4KG5 binding. Zwick *et al.* [2003] (antibody interactions)
- X5: The human Fab X5 was selected from a phage display library derived from an HIV-1 positive donor with a highly neutralizing serum it was selected for binding to purified gp120-CD4-coreceptor complexes the Fab neutralizes PBMC infection by a selection of HIV-1 primary isolates from clades A, B, C, D, E, F, and G, and neutralizes R5, X4, and R5X4 isolates it binds to a conserved epitope on gp120 induced by CD4 binding, its binding is slightly enhanced by CCR5 binding while CD4i MAb 17b binds the CCR5 binding site, X5 also competes with Fab b12 which overlaps with the CD4 binding site, suggesting the epitope for is near both the CD4 and CCR5 binding sites. Moulard *et al.* [2002] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 1127

**MAb ID** 8F101

HXB2 Location Env

**Author Location** gp120

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: • B clade HXB2 HIV component: gp120

Species (Isotype) mouse (IgG)

**Ab Type** gp120 CD4i, gp120-CD4 complex

**Research Contact** Ranajiit Pal, Advanced BioScience Lab, Inc. **References** Finnegan *et al.* 2002; Finnegan *et al.* 2001;

DeVico et al. 1995

**Keywords** antibody binding site definition and exposure, antibody generation, kinetics

8F101: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I and Cluster II MAbs required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but

not the CXCR4 co-receptor, binding to a fusion intermediate. 8F101 selectively stains gp120-CD4 complexes after dissociation from gp41, and did not stain cells arrested earlier than 30 min of co-culture, but 8F101 and cluster I and II MAbs co-localized at fusing cell interfaces at 30 min coculture. After extended co-culture, only 8F101 bound. Finnegan *et al.* [2002] (antibody binding site definition and exposure, kinetics)

- 8F101: Using a cell-fusion system, it was found CD4i antibodies 17b, 48d, and CG10 reacted faintly with Env expressing HeLA cells even in the absence of sCD4 or CD4 expressing target cells. Reactivity increased after sCD4 addition, but not after CD4 expressing target cell addition, and binding was not increased at the cell-to-cell CD4-Env interface. This suggests the CD4i co-receptor binding domain is largely blocked at the cell-fusion interface, and so CD4i antibodies would not be able access this site and neutralize cell-mediated viral entry. However, CD4i MAbs 8F101 and A32, that bind outside the co-receptor domain, had a different pattern. They reacted after the formation of gp120-CD4-CXCR4 tri-complexes, so co-receptor interactions allowed exposure of their epitopes. Finnegan *et al.* [2001] (antibody binding site definition and exposure)
- 8F101: MAbs specifically reactive to crosslinked gp120 and CD4 were derived (8F101, 8F102) conformation dependent competition studies indicate the epitope is immunogenic in infected humans. DeVico et al. [1995] (antibody generation)

**No.** 1128

MAb ID T22

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope

**Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

Ab Type Env oligomer

**Research Contact** P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

**References** Sugiura *et al.* 1999; Otteken *et al.* 1996; Earl *et al.* 1994

- T22: A comparison of 25 gp120 specific, conformation dependent MAbs was done T22 is part of a group of MAbs labeled AII all AII MAbs were broadly cross-reactive with gp160 from B-clade R5, X4, and R5X4 viruses, and could only partially blocked CD4 binding. Sugiura *et al.* [1999]
- T22: Pulse label experiments of 4 MAbs (D20, D27, T20, and T22) binding to noncleavable gp140 revealed that these anti-CD4BS MAbs bound with a delay, and that the epitope formed with a t 1/2 of about 10 minutes. Otteken *et al.* [1996]
- T22: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

No. 1129

MAb ID 2A2

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type N-term

References Weissenhorn et al. 1996

• Soluble gp41(21-166) forms a rod like structure that can be visualized with electron microscopy, and 2A2 binds to one end of the rod. Weissenhorn et al. [1996]

**No.** 1130

MAb ID AC4

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing yes

Immunogen vaccine

Vector/Type: protein HIV component:

gp160

Species (Isotype) mouse

Ab Type N-term

References Dickey et al. 2000

• AC4: Three MABs, ID6, AC4, and AD3 that bind to a discontinuous N-term first 204 aa of gp120 and generate ADCC were elicited through vaccination of BALBc mice with rec gp160 - these MAbs do not depend on glycosylation and are crossreactive with viruses from clades B and CRF01(AE). Dickey et al. [2000]

No. 1131

MAb ID AD3

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

**Neutralizing** yes

Immunogen vaccine

Vector/Type: protein HIV component:

gp160

Species (Isotype) mouse

Ab Type N-term

References Cook et al. 1994; Dickey et al. 2000

- AD3: There may be two Abs with this name that bind to the N-term region of gp120. Cook et al. [1994]; Dickey et al. [2000]
- AD3: Three MABs, ID6, AC4, and AD3 that bind to a discontinuous N-term first 204 aa of gp120 and generate ADCC were elicited through vaccination of BALBc mice with rec gp160 - these MAbs do not depend on glycosylation and are crossreactive with viruses from clades B and CRF01(AE). Dickey et al. [2000]

**No.** 1132

MAb ID AD3

**HXB2 Location** Env

Author Location gp120 (BH10)

**Epitope** 

**Neutralizing** 

**Immunogen** 

Species (Isotype) mouse (IgG1)

Ab Type N-term

References Dickey et al. 2000; Cook et al. 1994; Ugen et al. 1993

- AD3: NIH AIDS Research and Reference Reagent Program: 2342.
- AD3: There may be two Abs with this name that bind to the N-term region of gp120. Cook et al. [1994]; Dickey et al.
- AD3: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon – MAbs against the N-terminal half of gp120 do not inhibit gp120 binding to GalCer - binding of GalCer to gp120 does not inhibit MAb binding. Cook et al. [1994]

No. 1133

MAb ID ID6

**HXB2 Location** Env

Author Location gp120 (1-193 BH10)

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) mouse (IgG1)

Ab Type N-term

References Dickey et al. 2000; Cook et al. 1994; Ugen et al. 1993

- ID6: NIH AIDS Research and Reference Reagent Program:
- ID6: There may be two Abs with this name that bind to the N-term region of gp120. Cook et al. [1994]; Dickey et al. [2000]
- ID6: MAbs against the glycosphingolipid GalCer block HIV infection of normally susceptible CD4 negative cells from the brain and colon – MAbs against the N-terminal half of gp120 do not inhibit gp120 binding to GalCer - binding of GalCer to gp120 does not inhibit MAb binding. Cook et al. [1994]

No. 1134

MAb ID ID6

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing yes

Immunogen vaccine

*Vector/Type:* protein HIV component:

gp160

Species (Isotype) mouse (IgG2a)

**Ab Type** N-term

References Cook et al. 1994; Dickey et al. 2000

- ID6: There may be two Abs with this name that bind to the N-term region of gp120. Cook et al. [1994]; Dickey et al. [2000]
- · ID6: Three MABs, ID6, AC4, and AD3 that bind to a discontinuous N-term first 204 aa of gp120 and generate ADCC were elicited through vaccination of BALBc mice with rec gp160 - these MAbs do not depend on glycosylation and are crossreactive with viruses from clades B and CRF01(AE). Dickey et al. [2000]

No. 1135

MAb ID 11/68b

HXB2 Location Env

Author Location gp120

**Epitope** 

**Neutralizing** L (HXB2)

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

**Species (Isotype)** rat (IgG1)

**Ab Type** gp120 V1-V2

Research Contact Shotton and Dean

**References** Peet *et al.* 1998; Shotton *et al.* 1995; McKeating *et al.* 1993b

- 11/68b: 435 (Y/H) in C4 does not abrogate binding (John Moore, per comm, 1996).
- 11/68b: UK Medical Research Council AIDS reagent: ARP3041.
- 11/68b: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind 11/68b was not affected by V3 serine substitutions mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]
- 11/68b: Cross-competes with MAbs 62c, 66c, 66a, and CRA-4

   similar to MAb 62c HXB2 neutralization escape mutant had
   a D/N substitution at residue 185 non-reciprocal inhibition of
   binding of CRA-3 and CRA-6. Shotton *et al.* [1995]
- 11/68b: Changes at residues 183/184 (PI/SG) within V2, 435 (Y/H) in C4, abrogate binding. McKeating *et al.* [1993b]

**No.** 1136

MAb ID 62c

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG1)

Ab Type gp120 V1-V2

References Shotton et al. 1995

- 62c: UK Medical Research Council AIDS reagent: ARP3075.
- 62c: Cross-competes with MAbs 11/68b, 66c, 66a, and CRA-4 same cross-competition group as MAb 11/68b non-reciprocal inhibition of binding of CRA-3 and CRA-6 substitutions 176-177 FY/AT, 179-180 LD/DL, 183-184 PI/SG, and 191-193 YSL/GSS abrogate binding binds but does not neutralize Hx10. Shotton *et al.* [1995]

**No.** 1137

MAb ID CRA-6 (CRA6)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype) mouse

**Ab Type** gp120 V1-V2 **References** Shotton *et al.* 1995

 CRA-6: Called CRA6 – same competition group as CRA-3. Shotton et al. [1995]

**No.** 1138

MAb ID L15

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing P (weak)

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

**Ab Type** gp120 V1-V2

References Gorny & Zolla-Pazner 2004; Parren et al.

1997b; Ditzel et al. 1997

**Keywords** review, variant cross-recognition or cross-neutralization

- L15: In a review of Envelope binding MAbs in this database, V2-specific MAbs are noted to have some ability to neutralize HIV-1, but generally weak with limited cross-reactivity. L15 and L17 are Fabs specific for V2. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization, review)
- L15: gp120 immobilized on solid phase by capture with anti-CD4 BS MAb L72 was used for selection of Fabs 2 anti-V2 Fabs were obtained with very similar epitopes, L15 and L17 deletions in V1 and V2 abolished binding, and rodent anti-V2 MAbs SC258, CRA3, G3-G4,G3-136, BAT-085, and 52-684 all compete with L15. Ditzel *et al.* [1997]
- L15: Does not neutralize TCLA strains but neutralizes some primary isolates weakly. Parren *et al.* [1997b]

No. 1139

MAb ID T52

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope

Neutralizing

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

**Ab Type** gp120 V1-V2

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- T52: A comparison of 25 gp120 specific, conformation dependent MAbs was done T52 is one of two MAbs labeled B-II, that had limited cross-reactivity with seven clade B isolates and did not fully blocked CD4 binding deletion of V1/V2 loops abrogated binding. Sugiura *et al.* [1999]
- T52: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl *et al.* [1994]

**No.** 1140

MAb ID T54

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

Neutralizing no

Immunogen vaccine

HIV component: oligomeric gp140

Species (Isotype) mouse (IgG)

**Ab Type** gp120 V1-V2

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- T54: A comparison of 25 gp120 specific, conformation dependent MAbs was done - T54 is one of two MAbs labeled B-II, that had limited cross-reactivity with seven clade B isolates and did not fully blocked CD4 binding - deletion of V1/V2 loops abrogated binding. Sugiura et al. [1999]
- T54: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1141

MAb ID polyclonal

**HXB2 Location** Env **Author Location** Env

**Epitope** 

Neutralizing yes

Immunogen HIV-1 infection

Species (Isotype) human

**Ab Type** gp120 V1-V2 and V3-V5 References Gordon & Delwart 2000

• Primary isolates have great differences in susceptibility to neutralization - the variation in V1V2 and V3-V5 was measured by HTA in a set of viruses with a range of neutralization susceptibilities, and greater variability was uncorrelated with resistance to neutralization. Gordon & Delwart [2000]

**No.** 1142

**MAb ID** 1088

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

**Immunogen** 

Species (Isotype)

Ab Type gp120 V2

References Berman et al. 1997

• 1088: Binds weakly to 2/7 isolates from breakthrough cases from a MN gp120 vaccine trial. Berman et al. [1997]

**No.** 1143

**MAb ID** 110-B

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

HIV infected-cell lysate Vector/Type: Strain: B clade BRU HIV component:

HIV-1

Species (Isotype) mouse

Ab Type gp120 V2

Research Contact Hybridolabs, Institute Pasteur, Paris, France

References Moore et al. 1993a

Vector/Type: vaccinia Strain: B clade IIIB • 110-B: specific for BH10, does not bind to MN, RF, or SF-2 gp120 - binding inhibited by deletion of the V2 loop, and the following amino acid substitutions: 168 K/L, 176/177 FY/AT, 179/180 LD/DL, 183/184 PI/SG, and 192-194 YSL/GSS. Moore *et al.* [1993a]

**No.** 1144

**MAb ID** 1357

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 V2

Research Contact Susan (Zol-Zolla-Pazner

> las01@mcrcr6.med.nyu) (NYU Med.

Center)

References Gorny & Zolla-Pazner 2004; Ling et al. 2002;

Nyambi et al. 2000; Gorny et al. 2000;

Nyambi et al. 1998

Keywords antibody binding site definition and exposure, co-receptor, review

- 1357: This broad review of anti-Envelope MAbs notes that V2 MAbs are generally weakly neutralizing at best, and somewhat strain specific. Anti-V2 MAbs 1357, 1361, 1393 are nonneutralizing. Gorny & Zolla-Pazner [2004] (review)
- 1357: A pseudotyping assay showed that an X4 V3 loop peptide could enhance infectivity of X4 virus, R5 and R5X4 V3 loops peptides could enhance infectivity of an R5 virus, and R5X4 peptides could enhance infectivity of an R5X4 virus. Neither R5 nor R5X4 peptides influenced binding of CD4BS MAbs F105 and Ig1Gb12, but did increase binding of CD4i MAb 17b. Of three V2 MAbs, only 830A, not 2158 or 1357 was enhanced by V3 peptide binding. Ling et al. [2002] (antibody binding site definition and exposure, co-receptor)
- 1357: Blocks binding of MAb 697-D to rgp120, and doesn't react with a protein from which V1V2 has been deleted - binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared – no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer - V2 MAbs 697-D, 1357 and 1361 favored the monomer by approximately 2 fold. Gorny et al. [2000]
- 1357: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 anti-V2 MAbs, which showed weak and sporadic binding, with the most frequent binding to C and D clades. Nyambi et al. [2000]
- 1357: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 anti-V2 MAbs, which showed weak and sporadic binding, with the most frequent binding to C and D clades. Nyambi et al. [2000]
- 1357: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H - V2 Abs 697-D, 1361, and 1357 tended to bind very weakly with a similar pattern of specificity to virions, but bound well to soluble gp120: weak binding only to subtype D MAL. Nyambi et al. [1998]

**No.** 1145

MAb ID 1361
HXB2 Location Env
Author Location gp120
Epitope
Neutralizing
Immunogen vaccine

Vector/Type: protein HIV component:

gp120

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 V2

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

**References** Nyambi *et al.* 2000; Gorny *et al.* 2000; Nyambi *et al.* 1998

- 1361: Blocks binding of MAb 697-D to rgp120, and doesn't react with a protein from which V1V2 has been deleted binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, though anti-V3 and CD4BS MAbs reacted better with the oligomer and V2 and C5 tended to favor the monomer V2 MAbs 697-D, 1357 and 1361 favored the monomer by approximately 2 fold. Gorny *et al.* [2000]
- 1361: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 anti-V2 MAbs, which showed weak and sporadic binding, with the most frequent binding to C and D clades. Nyambi *et al.* [2000]
- 1361: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H V2 Abs 697-D, 1361, and 1357 tended to bind weakly with a similar pattern of specificity to virions, but bound well to soluble gp120: weak binding to 1/4 B clade viruses (CA5), and also weak binding to a subtype D virus, MAL. Nyambi *et al.* [1998]

**No.** 1146

**MAb ID** 1393A

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype)

Ab Type gp120 V2

References Gorny & Zolla-Pazner 2004; Nyambi et al.

2000

Keywords review, subtype comparisons

- 1393A: This broad review of anti-Envelope MAbs notes that V2
  MAbs are generally weakly neutralizing at best, and somewhat
  strain specific. Anti-V2 MAbs 1357, 1361, 1393A are nonneutralizing. Gorny & Zolla-Pazner [2004] (review)
- 1393A: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 anti-V2 MAbs, which showed weak and sporadic binding, with the most frequent binding to C and D clades. Nyambi *et al.* [2000] (subtype comparisons)

**No.** 1147

**MAb ID** 2158

**HXB2 Location** Env

Author Location gp120 (LAI)

**Epitope** 

Subtype B

Neutralizing

**Immunogen** 

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 V2

Research Contact Susan Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu) (NYU Med.

Center)

References Pinter et al. 2004; Ling et al. 2004; Ling et al.

2002

Keywords antibody binding site definition and exposure,

co-receptor, variant cross-recognition or cross-

neutralization

- 2158: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of 17b and F105 was decreased by trypsin, but increased by thrombin. V2 MAbs 830A and 2158 were decreased by trypsin, uneffected by thrombin. Thrombin may increase HIV-induced cell fusion in blood by causing a conformational activating shift in gp120. Ling et al. [2004] (antibody binding site definition and exposure)
- 2158: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Three anti-V2 MAb were tested both 2158 and 830A bound more strongly to JRFL, but neutralized SF162, and not neutralize JRFL. Thus V2 domains are better neutralization targets in SF162. Pinter et al. [2004] (variant cross-recognition or cross-neutralization)
- 2158: A pseudotyping assay showed that an X4 V3 loop peptide could enhance infectivity of X4 virus, R5 and R5X4 V3 loops peptides could enhance infectivity of an R5 virus, and R5X4 peptides could enhance infectivity of an R5X4 virus. Neither R5 nor R5X4 peptides influenced binding of CD4BS MAbs F105 and Ig1Gb12, but did increase binding of CD4i MAb 17b. Of three V2 MAbs, only 830A, not 2158 or 1357 was enhanced by V3 peptide binding. Ling et al. [2002] (antibody binding site definition and exposure, co-receptor)

**No.** 1148

MAb ID 66a

**HXB2 Location** Env

**Author Location** gp120

Epitope

Neutralizing L (HXB2)

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) mouse (IgG1)

**Ab Type** gp120 V2

References Shotton et al. 1995

• 66a: UK Medical Research Council AIDS reagent: ARP3074.

• 66a: Substitutions 176-177 FY/AT, 179-180 LD/DL, 183-184 PI/SG, and 191-193 YSL/GSS abrogate binding - same competition group as CRA4. Shotton et al. [1995]

No. 1149

MAb ID 66c

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L (HXB2)

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 V2

References Shotton et al. 1995

• 66c: Substitutions 176-177 FY/AT, 179-180 LD/DL, 183-184 PI/SG, and 191-193 YSL/GSS abrogate binding - same competition group as CRA4. Shotton et al. [1995]

No. 1150

MAb ID 684-238 (52-684-238, 52-684)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 V2

Research Contact Gerry Robey, Abbott Laboratories

References Ditzel et al. 1997; Moore & Sodroski 1996; Ditzel et al. 1995; Gorny et al. 1994; Thali et al. 1993; Moore et al. 1993a

- 684-238: Limited reciprocal enhancement of binding with anti-V3 and C4 region antibodies – reciprocal inhibition with V2 region antibodies. Moore & Sodroski [1996]
- 684-238: Does not compete with IgG1b12, reciprocal inhibition with MAbs L39, L40, and L78. Ditzel et al. [1995]
- 684-238: Weakly neutralizing, IC 50 = 84 mug/ml. Gorny et al.
- 684-238: Specific for BH10 or HXB2, does not bind to MN, RF, or SF-2 gp120 – neutralizes BH10 – binding inhibited by deletion of the V2 loop, and the following amino acid substitutions: 176/177FY/AT, 179/180LD/DL, 183/184PI/SG, and 192-194YSL/GSS. Moore et al. [1993a]

No. 1151

**MAb ID** 830A

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype)

Ab Type gp120 V2

Research Contact Susan Zolla-Pazner

References Gorny et al. 2005; Pinter et al. 2004; Ling et al. 2004; Gorny & Zolla-Pazner 2004; Ling

et al. 2002; Nyambi et al. 2000

Keywords antibody binding site definition and exposure, co-receptor, review, subtype comparisons, variant cross-recognition or crossneutralization

- 830A: 2909 is a human anti-Env NAb that was selected by neutralization assay and binds to the quaternary structure on the intact virion. ELISA-based competition assays and subsequent mutational analysis determined that the CD4BS and V2 and V3 loops contribute to the 2909 epitope: 2909 binding was inhibited by MAbs 447-52d (anti-V3), 830A (anti-V2), and IgG1b12 (anti-CD4BS) and sCD4. 2909 was not inhibited by MAbs 670, 1418, nor 2G12. Gorny et al. [2005]
- 830A: This broad review of anti-Envelope MAbs notes that V2 MAbs are generally weakly neutralizing at best, and somewhat strain specific. 830A neutralizes SF162. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization,
- 830A: The role of serine proteases on HIV infection was explored. Trypsin decreased the binding of most Env MAb tested and diminished cell fusion of H9 cells infected with HIV-1 LAI virus (H9/IIIB) to MAGI cells. In contrast, thrombin increased the binding of MAbs to gp120 epitopes near the CD4 and CCR5 binding sites, and increased cell fusion. Binding of 17b and F105 was decreased by trypsin, but increased by thrombin. V2 MAbs 830A and 2158 were decreased by trypsin, uneffected by thrombin. Thrombin may increase HIV-induced cell fusion in blood by causing a conformational activating shift in gp120. Ling et al. [2004] (antibody binding site definition and exposure)
- 830A: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neutralization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. Three anti-V2 MAb were tested - both 2158 and 830A bound more strongly to JRFL, but neutralized SF162, and did not neutralize JRFL. Thus V2 domains are better neutralization targets in SF162. Pinter et al. [2004] (variant cross-recognition or cross-neutralization)
- 830A: A pseudotyping assay showed that an X4 V3 loop peptide could enhance infectivity of X4 virus, R5 and R5X4 V3 loops peptides could enhance infectivity of an R5 virus, and R5X4 peptides could enhance infectivity of an R5X4 virus. Neither R5 nor R5X4 peptides influenced binding of CD4BS MAbs F105 and Ig1Gb12, but did increase binding of CD4i MAb 17b. Of three V2 MAbs, only 830A, not 2158 or 1357 was enhanced by V3 peptide binding. Ling et al. [2002] (antibody binding site definition and exposure, co-receptor)
- 830A: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 anti-V2 MAbs, which showed weak and sporadic binding, with the most frequent

recognition or cross-neutralization, subtype comparisons)

No. 1152

MAb ID CRA-3 (CRA3)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) mouse (IgG2a)

Ab Type gp120 V2

Research Contact Mark Page, NIBSC AIDS reagent project, Pot-

ters Bar, Herts, UK

References Ditzel et al. 1997; Moore & Sodroski 1996;

Shotton et al. 1995; Thali et al. 1993; Moore

et al. 1993a; Moore & Ho 1993

• CRA-3: UK Medical Research Council AIDS reagent: ARP324.

- CRA-3: Many MAbs enhance binding, including some anti-C5, C1, V4, and C4 MAbs - enhances binding of only a small number of anti-V3 loop MAbs. Moore & Sodroski [1996]
- CRA-3: Called CRA3 Same competition group as CRA6. Shotton *et al.* [1995]
- CRA-3: Conformational, does not bind well to denatured gp120. Moore & Ho [1993]
- CRA-3: specific for BH10 or HXB2, does not bind to MN, RF, or SF-2 gp120 - binding inhibited by deletion of the V2 loop, and the following amino acid substitutions: 176/177 FY/AT, 179/180 LD/DL, 183/184 PI/SG, and 192-194 YSL/GSS - epitope probably involves stem of V1/V2 loop structure. Moore et al. [1993a]

**No.** 1153

MAb ID CRA-4 (CRA4)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L (HXB2)

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120 V2

Research Contact Mark Page, NIBS, MRC AIDS reagent repos-

itory, ARP 325

References Moore & Sodroski 1996; Shotton et al. 1995; Thali et al. 1993; Moore et al. 1993a; Moore

& Ho 1993; McKeating et al. 1993b

- CRA-4: UK Medical Research Council AIDS reagent: ARP325.
- CRA-4: The only MAbs that enhanced binding were anti-V3 MAb 5G11 and anti-C1 MAb 135/9 binding – reciprocal inhibition of anti-V2 MAbs. Moore & Sodroski [1996]
- CRA-4: Cross-competes with MAbs 11/68b, 62c, 66c, 66a similar to 66c and 66a - non-reciprocal inhibition by MAbs 12b, 60b and CRA-6. Shotton et al. [1995]

binding to C and D clades. Nyambi et al. [2000] (variant cross-V2, 435 (Y/H) in C4, abrogate binding – type-specific neutralization. McKeating et al. [1993b]

- CRA-4: Conformational, does not bind well to denatured gp120. Moore & Ho [1993]
- CRA-4: Specific for BH10 and HXB2, does not bind to MN, RF, or SF-2 gp120 - binding inhibited by deletion of the V2 loop, and the following amino acid substitutions: 176/177 FY/AT, 179/180 LD/DL, 183/184 PI/SG, and 192-194 YSL/GSS. Moore et al. [1993a]

No. 1154

MAb ID L17

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** 

**Immunogen** 

Species (Isotype) human

Ab Type gp120 V2

References Gorny & Zolla-Pazner 2004; Kwong et al.

2002; Parren et al. 1998a; Ditzel et al. 1997

Keywords antibody binding site definition and exposure, binding affinity, review, variant crossrecognition or cross-neutralization

- L17: In a review of Envelope binding MAbs in this database, V2-specific MAbs are noted to have some ability to neutralize HIV-1, but generally weakly with limited cross-reactivity. L15 and L17 are Fabs specific for V2. Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, review)
- L17: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-2999) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong et al. [2002] (antibody binding site definition and exposure)
- L17: The rank order of Fab binding affinity to monomeric gp120 (Loop 2 > 3B3 > b12 = DO8i > b11 > b3 > b14 > b13> DO142-10 > DA48 > L17) was markedly different than Fab binding affinity to the mature oligomeric form (3B3 > b12 > DO142-10 > Loop 2 > b11 > L17 > b6 > DO8i > b14 > DA48 > b3 > b13) and binding to oligomeric form and neutralization were correlated for both Fabs and MAbs - authors suggest

that neutralization is determined by the fraction of Ab sites occupied on a virion irrespective of the epitope. Parren et al. [1998a] (binding affinity)

**No.** 1155

MAb ID SC258 (52-581-SC258)

**HXB2 Location** Env Author Location gp120 **Epitope** 

**Neutralizing** L Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: gp120

Species (Isotype) mouse

Ab Type gp120 V2

Research Contact Gerry Robey, Abbott Laboratories

References He et al. 2002; Ditzel et al. 1997; Trkola et al. 1996a; Moore & Sodroski 1996; Ditzel et al. 1995; Moore et al. 1994b; Yoshiyama et al. 1994; Gorny et al. 1994; Thali et al. 1993; Moore et al. 1993a

- SC258: Transgenic mice carrying human genes allowing production of fully human MAbs were used to rapidly create a panel of anti-HIV gp120 MAb producing hybridomas by immunization with HIV SF162 gp120 - the previously described human MAbs 5145A(CD4BS), 4117C (plus others, V3) and 697D (and SC258, V2) were used as controls. He et al. [2002]
- SC258: Several MAbs binding to various gp120 epitopes enhance binding, but the only MAb that SC258 enhanced binding of was anti-CD4 binding site MAb F91 - reciprocal inhibition with V2 region antibodies. Moore & Sodroski [1996]
- SC258: Does not inhibit gp120 interaction with CCR-5 in a MIP-1beta-CCR-5 competition study - listed as not neutralizing. Trkola et al. [1996a]
- SC258: Does not compete with IgG1b12 reciprocal inhibition with MAbs L39, L40, and L78. Ditzel et al. [1995]
- SC258: Very poor reactivity with gp120 molecules outside of clade B. Moore et al. [1994b]
- SC258: HIV-1 RF V2 substitutions 177 Y/H and 179 L/P in the V2 loop of RF reduce affinity – 177 Y/H inhibits SC258 neutralization. Yoshiyama et al. [1994]
- SC258: Called 52-581-SC258 binds to BH10, MN, and RF gp120 – neutralizes BH10 – binding inhibited by deletion of the V2 loop, and the following amino acid substitutions: 176/177 FY/AT, 179/180 LD/DL, 183/184 PI/SG, and 192-194 YSL/GSS. Moore et al. [1993a]

**No.** 1156

MAb ID L25

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L (weak)

Immunogen HIV-1 infection

Species (Isotype) human (IgG1)

**Ab Type** gp120 V2-CD4BS References Gorny & Zolla-Pazner 2004; Parren et al. Author Location gp120

1997b; Ditzel et al. 1997; Ditzel et al. 1995

**Keywords** antibody binding site definition and exposure, review, variant cross-recognition or crossneutralization

- L25: In a review of Envelope binding MAbs in this database, V2-specific MAbs are noted to have some ability to neutralize HIV-1, but generally weakly with limited cross-reactivity. L25, L39, L40 and L78 are Fabs specific for a complex binding site involving V2 and elements of the sCD4 binding site regions; among these only L25 and L78 mediate weak neutralization of some TCLA strains. Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, review)
- L25: gp120 immobilized on solid phase by capture with anti-CD4 BS MAb L72 was used for selection of Fabs – a single anti-V2-CD4 BS Fab was obtained with with sensitivity to substitutions in the V2 and CD4 BS regions - rodent anti-V2 MAb SC258 competes with L25. Ditzel et al. [1997]
- L25: Neutralizes TCLA strains weakly, but not primary isolates. Parren et al. [1997b]

No. 1157

MAb ID L39

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type gp120 V2-CD4BS

References Gorny & Zolla-Pazner 2004; Ditzel et al. 1995 Keywords antibody binding site definition and exposure, review, variant cross-recognition or crossneutralization

- L39: In a review of Envelope binding MAbs in this database, V2-specific MAbs in are noted have some ability to neutralize HIV-1, but generally weakly with limited cross-reactivity. L25, L39, L40 and L78 are Fabs specific for a complex binding site involving V2 and elements of the sCD4 binding site regions; among these only L25 and L78 mediate weak neutralization of some TCLA strains. Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, review)
- L39: This Fab does not inhibit sCD4 binding, but is inhibited by sCD4, probably due to conformational changes – it is competed by anti-V2 MAbs, and sensitive to amino acid substitutions in the V3 loop (similar patterns were observed for L39 and L78 gp120 amino acid substitutions enhancing or reducing binding) - does not compete with CD4BS MAbs, but is sensitive to amino acid changes at positions 368 and 370 - binding unaffected by deglycosylation - reciprocal inhibition with V2 MAbs SC258 and 684-238 - heavy and light chain variable region sequence is available. Ditzel et al. [1995]

No. 1158

MAb ID L40

**HXB2 Location** Env

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp120 V2-CD4BS

**References** Gorny & Zolla-Pazner 2004; Ditzel *et al.* 1995 **Keywords** antibody binding site definition and exposure, responses in children, variant cross-recognition or cross-neutralization

- L40: In a review of Envelope binding MAbs in this database, V2-specific MAbs are noted to have some ability to neutralize HIV-1, but generally weakly with limited cross-reactivity. L25, L39, L40 and L78 are Fabs specific for a complex binding site involving V2 and elements of the sCD4 binding site regions; among these only L25 and L78 mediate weak neutralization of some TCLA strains. Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, responses in children)
- L40: This Fab does not inhibit sCD4 binding, but is inhibited by sCD4, probably due to conformational changes it is competed by anti-V2 MAbs, and sensitive to amino acid substitutions in the V3 loop (similar patterns were observed for L40 and L78 gp120 amino acid substitutions enhancing or reducing binding) does not compete with CD4BS MAbs, but is sensitive to amino acid changes at positions 368 and 370 binding only partially affected by deglycosylation reciprocal inhibition with V2 MAbs SC258 and 684-238 heavy and light chain variable region sequence is available. Ditzel et al. [1995]

**No.** 1159

MAb ID L78

**HXB2 Location** Env

Author Location gp120

Epitope

**Neutralizing** L

Immunogen HIV-1 infection Species (Isotype) human (IgG1 $\kappa$ ) Ab Type gp120 V2-CD4BS

References Gorny & Zolla-Pazner 2004; Kwong et al.

2002; Ditzel et al. 1995

**Keywords** antibody binding site definition and exposure, antibody sequence, variable domain, review, variant cross-recognition or cross-neutralization

- L78: In a review of Envelope binding MAbs in this database, V2-specific MAbs are noted to have some ability to neutralize HIV-1, but generally weakly with limited cross-reactivity. L25, L39, L40 and L78 are Fabs specific for V2 that are also associated with sCD4 binding site regions; among these only L25 and L78 mediate weak neutralization of some TCLA strains. Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, review)
- L78: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS

neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-2999) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong *et al.* [2002] (antibody binding site definition and exposure)

• L78: Substitutions at V2: (152/153 GE/SM, 183/184 PI/SG, 191/193 YL/GS), 262 N/T, V3 (314 G/W), CD4BS (257 T/R, 368 D/R, 370 E/R) inhibit binding, and some C4 and C5 substitutions enhance binding – this Fab does not inhibit sCD4 binding, but is inhibited by sCD4, probably due to conformational changes – it is competed by anti-V2 MAbs, and sensitive to amino acid substitutions in the V3 loop – does not compete with CD4BS MAbs, but is sensitive to amino acid changes at positions 368 and 370 – Fab neutralizes MN and LAI – binding unaffected by deglycosylation – reciprocal inhibition with V2 MAbs SC258 and 684-238 – heavy and light chain variable region sequence is available. Ditzel et al. [1995] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, antibody sequence, variable domain)

**No.** 1160

MAb ID

HXB2 Location Env

**Author Location** gp120

**Epitope** 

Subtype A, B, C

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Gilljam et al. 1999

Sera from individuals with infections of HIV-1 subtype A-E were tested against purified proteins from primary PBMC cultures. Sera reactivity tended not to be strongly related to subtype, rather probably reflected the sum of reactivities to conserved and variable regions in the proteins. V3 peptide comparisons showed some preference for within subtype binding. Gilljam et al. [1999]

No. 1161

**MAb ID** 10D8

**HXB2 Location** Env

**Author Location** gp160 (V3) (303–338)

**Epitope** 

Subtype B

Neutralizing

Immunogen

Species (Isotype) human

**Ab Type** gp120 V3

References Callahan et al. 1991

• 10D8: Polyanionic polysaccharides were proposed to inhibit viral functions such as binding and syncytia formation through interations mediated through the local high positive charge density in the V3 loop. The binding of this anti-V3 antibody is inhibited by dextransulfate. Callahan *et al.* [1991]

No. 1162
MAb ID 10F6
HXB2 Location Env
Author Location gp160 (V3) (303–338)
Epitope
Subtype B
Neutralizing
Immunogen
Species (Isotype) human
Ab Type gp120 V3
References Callahan et al. 1991

• 10F6: Polyanionic polysaccharides were proposed to inhibit viral functions such as binding and syncytia formation through interations mediated through the local high positive charge density in the V3 loop. The binding of this anti-V3 antibody is inhibited by dextransulfate. Callahan *et al.* [1991]

No. 1163
MAb ID 110.J
HXB2 Location Env
Author Location gp120
Epitope
Neutralizing
Immunogen
Species (Isotype)
Ab Type gp120 V3

**Research Contact** F. Traincard, Pasteur Institute, France

References Moore & Sodroski 1996; Thali et al. 1993

110.J: Binds to carboxy-terminal side of the V3 loop – reciprocal binding inhibition with other anti-V3 and anti-C4 MAbs – and reciprocal enhanced binding of some anti-V2 MAbs and anti-CD4 binding site MAbs. Moore & Sodroski [1996]

• 110.J: Inhibits sCD4-inducible anti-CD4 binding site MAb 48d. Thali *et al.* [1993]

No. 1164 MAb ID 11G5 HXB2 Location Env Author Location gp160 (V3) (303–338) Epitope Subtype B Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 V3

References Callahan et al. 1991

• 11G5: Polyanionic polysaccharides were proposed to inhibit viral functions such as binding and syncytia formation through interations mediated through the local high positive charge density in the V3 loop. The binding of this anti-V3 antibody is inhibited by dextransulfate. Callahan *et al.* [1991]

No. 1165 MAb ID 2182 HXB2 Location Env Author Location (JRCSF) Epitope

Subtype B Neutralizing P

 $\begin{array}{c} \textbf{Immunogen} & \text{HIV-1 infection} \\ \textbf{Species} & \textbf{(Isotype)} & \text{human} & \textbf{(IgG1}\lambda) \end{array}$ 

**Ab Type** gp120 V3 **Research Contact** Susan

earch Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

References Pinter et al. 2004; Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Gorny et al. 2002

**Keywords** antibody binding site definition and exposure, antibody generation, review, subtype comparisons, variant cross-recognition or cross-neutralization

- 2182: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. The set that can cross-neutralize primary isolates (2182, 2191, 2219, 2412, 2442, 2456) bind V3 but are conformationally sensitive, suggesting some structural conservation despite sequence variation. These MAbs have distinct epitopes relative to 447-52D, a MAb directed at the tip of the V3 loop that also can neutralize many primary isolates. Inter-clade cross-neutralization by these anti-V3 MAbs is reduced. Gorny & Zolla-Pazner [2004] (review, subtype comparisons)
- 2182: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using a JR-CSF fusion protein, and could neutralize 6/13 B clade viruses. Gorny et al. [2004] (antibody binding site definition and exposure)
- 2182: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. 5/6 anti-V3 MAbs had similar binding affinity to soluble SF162 and JR-FL rgp120s, although the V3 loop differs at three positions (HigpgrafyTtgE for JR-FL and TigpgrafyAtgD for SF162). Only the V3 MAb that had a different affinity was 2182, which bound to JRFL with higher affinity. Even 2182 preferentially neutralized SF162, however, the JRFL gp120 backbone with the SF162 V1V2 region was the more neutralization sensitive than pure SF162. Pinter et al. [2004] (variant cross-recognition or cross-neutralization)
- 2182: Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated from cells of asymptomatic HIV-1-infected individuals by selection of heterhybridomas using a V3-fusion protein (V3-fp), the HIV-1

JRCSF V3 loop inserted into a truncated murine leukemia virus gp70 – the six new MAbs all bind to the tip of the V3 loop and cross-compete with the MAb 447-52D and are conformationally sensitive - MAbs showed cross-clade binding to native, intact virions of clades A(N=2), B(N=4), and F(N=2), limited binding to C(N=3) and D(N=3), and did not bind to CRF01(subtype E, N=2) – the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay - five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization studies), 654 (anti-CD4BS used as a conformationsensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control), MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades) – 5/6 MAbs were derived from individuals infected in the US, presumably with clade B, and one, 2182, was derived from an individual who was infected abroad with clade A who is presently living in New York city – 2412 and 2456 were produced from cells obtained from the same individual, while the other MAbs were each generated from different subjects – 2182 bound to 8/16 of the diverse isolates, not to any clade C or CRF01. Gorny et al. [2002] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

No. 1166
MAb ID 2191
HXB2 Location Env
Author Location (JRCSF)
Epitope
Subtype B
Neutralizing P
Immunogen HIV-1 infection
Species (Isotype) human (IgG1λ)
Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

**References** Pinter *et al.* 2004; Gorny *et al.* 2004; Gorny & Zolla-Pazner 2004; Gorny *et al.* 2002

**Keywords** antibody binding site definition and exposure, antibody generation, review, subtype comparisons, variant cross-recognition or cross-neutralization

• 2191: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. The set that can cross-neutralize primary isolates (2182, 2191, 2219, 2412, 2442, 2456) bind V3 but are conformationally sensitive, suggesting some structural conservation despite sequence variation. These MAbs have distinct epitopes relative to 447-52D, a MAb directed at the tip of the V3 loop that also can neutralize many primary isolates. Interclade cross-neutralization by these anti-V3 MAbs is reduced. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization, review, subtype comparisons)

 2191: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using a JR-CSF fusion protein, and could neutralize 8/13 B clade viruses. Gorny *et al.* [2004] (antibody binding site definition and exposure)

- 2191: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. 5/6 anti-V3 MAbs, including 2191, had similar binding affinity to soluble SF162 and JR-FL rgp120s, although the V3 loop differs at three positions (HigpgrafyTtgE for JR-FL and TigpgrafyAtgD for SF162). Pinter et al. [2004] (variant cross-recognition or cross-neutralization)
- 2191: Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated from cells of asymptomatic HIV-1-infected individuals by selection of heterhybridomas using a V3-fusion protein (V3-fp), the HIV-1 JRCSF V3 loop inserted into a truncated murine leukemia virus gp70 - the six new MAbs all bind to the tip of the V3 loop and cross-compete with the MAb 447-52D and are conformationally sensitive - MAbs showed cross-clade binding to native, intact virions of clades A(N=2), B(N=4), and F(N=2), limited binding to C(N=3) and D(N=3), and did not bind to CRF01(subtype E, N=2) - the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay – five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization studies), 654 (anti-CD4BS used as a conformation-sensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control), MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades) – 5/6 MAbs were derived from individuals infected in the US, presumably with clade B, and one, 2182, was derived from an individual who was infected abroad with clade A who is presently living in New York city - 2412 and 2456 were produced from cells obtained from the same individual, while the other MAbs were each generated from different subjects - 2191 bound to 10/16 of the diverse isolates, not to any clade D or CRF01. Gorny et al. [2002] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

No. 1167
MAb ID 2219
HXB2 Location Env
Author Location (JRCSF)
Epitope
Subtype B
Neutralizing P
Immunogen HIV-1 infection
Species (Isotype) human (IgG1λ)
Ab Type gp120 V3

Env Antibodies Tables

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med. Center)

References Pinter et al. 2005; Pinter et al. 2004; Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Gorny et al. 2002

**Keywords** antibody binding site definition and exposure, antibody generation, review, subtype comparisons, variant cross-recognition or cross-neutralization

- 2219: This study is about the V2 MAb C108g, that is type-specific and neutralizes BaL and HXB2. JR-FL is a neutralization resistant strain; modification of JRFL at V2 positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The modification in V2 also increased neutralization sensitivity to V3 MABs 4117c, 2219, 2191, and 447-52D, but only had minor effects on neutralization by CD4BS MAb 5145A, and broadly neutralizing MAbs IgG1b12, 2G12, and 2F5. Binding to CCR5 was completely inhibited by two V3 MAbs, 4117C and 2219, and was substantially inhibited by 2G12, but was not inhibited by C108g. Pinter et al. [2005] (antibody binding site definition and exposure)
- 2219: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. The set that can cross-neutralize primary isolates (2182, 2191, 2219, 2412, 2442, 2456) bind V3 but are conformationally sensitive, suggesting some structural conservation despite sequence variation. These MAbs have distinct epitopes relative to 447-52D, a MAb directed at the tip of the V3 loop that also can neutralize many primary isolates. Interclade cross-neutralization by these anti-V3 MAbs is reduced. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization, review, subtype comparisons)
- 2219: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4-induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using a JR-CSF fusion protein, and could neutralize 6/13 B clade viruses. Gorny et al. [2004] (antibody binding site definition and exposure)
- 2219: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. 5/6 anti-V3 MAbs, including 2219, had similar binding affinity to soluble SF162 and JR-FL rgp120s, although the V3 loop differs at three positions (HigpgrafyTtgE for JR-FL and TigpgrafyAtgD for SF162). Pinter et al. [2004] (variant cross-recognition or cross-neutralization)

(Zol- • 2219: Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated from cells of asymptomatic HIV-1-infected individuals by selection of heterhybridomas using a V3-fusion protein (V3-fp), the HIV-1 JRCSF V3 loop inserted into a truncated murine leukemia virus gp70 – the six new MAbs all bind to the tip of the V3 loop and cross-compete with the MAb 447-52D and are conformationally sensitive - MAbs showed cross-clade binding to native, intact virions of clades A(N=2), B(N=4), and F(N=2), limited binding to C(N=3) and D(N=3), and did not bind to CRF01(subtype E, N=2) - the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay - five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization studies), 654 (anti-CD4BS used as a conformation-sensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control), MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades) – 5/6 MAbs were derived from individuals infected in the US, presumably with clade B, and one, 2182, was derived from an individual who was infected abroad with clade A who is presently living in New York city - 2412 and 2456 were produced from cells obtained from the same individual, while the other MAbs were each generated from different subjects – 2219 bound to 13/16 of the diverse isolates. Gorny et al. [2002] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

No. 1168
MAb ID 2412
HXB2 Location Env
Author Location gp120 (V3) (JRCSF)
Epitope
Subtype B
Neutralizing P
Immunogen HIV-1 infection
Species (Isotype) human (IgG1\(\lambda\))
Ab Type gp120 V3

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med. Center)

**References** Gorny *et al.* 2004; Gorny & Zolla-Pazner 2004; Gorny *et al.* 2002

**Keywords** antibody binding site definition and exposure, antibody generation, review, subtype comparisons, variant cross-recognition or cross-neutralization

• 2412: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. The set that can cross-neutralize primary isolates (2182, 2191, 2219, 2412, 2442, 2456) bind V3 but are conformationally sensitiive, suggesting some structural conservation despite sequence variation. These MAbs have distinct epitopes relative to 447-52D, a MAb directed at the tip of the V3 loop that also can neutralize many primary isolates. Inter-clade cross-neutralization by these anti-V3 MAbs is reduced. Gorny & Zolla-Pazner [2004] (antibody binding site definition and exposure, review)

- 2412: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using a JR-CSF fusion protein, and could neutralize 4/13 B clade viruses. Gorny et al. [2004] (antibody binding site definition and exposure)
- 2412: Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated from cells of asymptomatic HIV-1-infected individuals by selection of heterhybridomas using a V3-fusion protein (V3-fp), the HIV-1 JRCSF V3 loop inserted into a truncated murine leukemia virus gp70 - the six new MAbs all bind to the tip of the V3 loop and cross-compete with the MAb 447-52D and are conformationally sensitive - MAbs showed cross-clade binding to native, intact virions of clades A(N=2), B(N=4), and F(N=2), limited binding to C(N=3) and D(N=3), and did not bind to CRF01(subtype E, N=2) – the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay - five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization studies), 654 (anti-CD4BS used as a conformation-sensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control), MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades) – 5/6 MAbs were derived from individuals infected in the US, presumably with clade B, and one, 2182, was derived from an individual who was infected abroad with clade A who is presently living in New York city – 2412 and 2456 were produced from cells obtained from the same individual, while the other MAbs were each generated from different subjects – 2412 bound to 7/16 of the diverse isolates, and did not bind to any of the clade C, D or CRF01 viruses. Gorny et al. [2002] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

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No. 1169
        MAb ID 2442
 HXB2 Location Env
 Author Location (JRCSF)
         Epitope
        Subtype B
    Neutralizing P
    Immunogen HIV-1 infection
Species (Isotype) human (IgG1\lambda)
        Ab Type gp120 V3
Research Contact Susan
                               Zolla-Pazner
                                                    (Zol-
                 las01@mcrcr6.med.nyu)
                                          (NYU
      References Gorny et al. 2004; Gorny & Zolla-Pazner
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2004; Gorny et al. 2002

**Keywords** antibody binding site definition and exposure, antibody generation, review, variant crossrecognition or cross-neutralization

• 2442: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary

isolates. The set that can cross-neutralize primary isolates (2182, 2191, 2219, 2412, 2442, 2456) bind V3 but are conformationally sensitive, suggesting some structural conservation despite sequence variation. These MAbs have distinct epitopes relative to 447-52D, a MAb directed at the tip of the V3 loop that also can neutralize many primary isolates. Interclade cross-neutralization by these anti-V3 MAbs is reduced. Gorny & Zolla-Pazner [2004] (variant cross-recognition or cross-neutralization, review)

- 2442: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using a JR-CSF fusion protein, and could neutralize 9/13 B clade viruses. Gorny et al. [2004] (antibody binding site definition and exposure)
- 2442: Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated from cells of asymptomatic HIV-1-infected individuals by selection of heterhybridomas using a V3-fusion protein (V3-fp), the HIV-1 JRCSF V3 loop inserted into a truncated murine leukemia virus gp70 - the six new MAbs all bind to the tip of the V3 loop and cross-compete with the MAb 447-52D and are conformationally sensitive - MAbs showed cross-clade binding to native, intact virions of clades A(N=2), B(N=4), and F(N=2), limited binding to C(N=3) and D(N=3), and did not bind to CRF01(subtype E, N=2) - the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay - five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization studies), 654 (anti-CD4BS used as a conformation-sensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control), MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades) – 5/6 MAbs were derived from individuals infected in the US, presumably with clade B, and one, 2182, was derived from an individual who was infected abroad with clade A who is presently living in New York city – 2412 and 2456 were produced from cells obtained from the same individual, while the other MAbs were each generated from different subjects - 2442 bound to 13/16 of the diverse isolates. Gorny et al. [2002] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, review)

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No. 1170
        MAb ID 2456
 HXB2 Location Env
Author Location (JRCSF)
        Epitope
        Subtype B
    Neutralizing P
    Immunogen HIV-1 infection
Species (Isotype) human (IgG1\lambda)
        Ab Type gp120 V3
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Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) Med. Center)

2004: Gornv et al. 2002

Keywords antibody binding site definition and exposure, review

- 2456: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. The set that can cross-neutralize primary isolates (2182, 2191, 2219, 2412, 2442, 2456) bind V3 but are conformationally senstitive, suggesting some structural conservation despite sequence variation. These MAbs have distinct epitopes relative to 447-52D, a MAb directed at the tip of the V3 loop that also can neutralize many primary isolates. Inter-clade cross-neutralization by these anti-V3 MAbs is reduced. Gorny & Zolla-Pazner [2004] (review)
- 2456: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using a JR-CSF fusion protein, and could neutralize 4/12 B clade viruses. Gorny et al. [2004] (antibody binding site definition and exposure)
- 2456: Conformation-dependent anti-V3 loop Abs may be more cross-reactive, so six new V3 MAbs were generated from cells of asymptomatic HIV-1-infected individuals by selection of heterhybridomas using a V3-fusion protein (V3-fp), the HIV-1 JRCSF V3 loop inserted into a truncated murine leukemia virus gp70 - the six new MAbs all bind to the tip of the V3 loop and cross-compete with the MAb 447-52D and are conformationally sensitive - MAbs showed cross-clade binding to native, intact virions of clades A(N=2), B(N=4), and F(N=2), limited binding to C(N=3) and D(N=3), and did not bind to CRF01(subtype E, N=2) - the strength binding was highly correlated with percent neutralization using the ghost cell or PHA blast assay - five well-characterized MAbs were used as controls: anti-V3 447-52D (anti-V3 MAb for competition and neutralization studies), 654 (anti-CD4BS used as a conformation-sensitive MAb control), 1331A (anti-C5 used as a linear binding site MAb control), MAb 246 (anti-gp41 MAb that bound to primary isolates of all clades) – 5/6 MAbs were derived from individuals infected in the US, presumably with clade B, and one, 2182, was derived from an individual who was infected abroad with clade A who is presently living in New York city – 2412 and 2456 were produced from cells obtained from the same individual, while the other MAbs were each generated from different subjects – 2456 bound to 12/16 of the diverse isolates. Gorny et al. [2002]

No. 1171 **MAb ID** 2483 **HXB2 Location** Env **Author Location** Env (JR-CSF) **Epitope** Subtype B Neutralizing P **Immunogen** Species (Isotype) human Ab Type gp120 V3

References Gorny et al. 2004; Gorny & Zolla-Pazner Research Contact Dr. Zolla-Pazner, Veterans Affairs Center, NY, NY. zollas01@endeavor.med.nvu.edu

References Gorny et al. 2004

**Keywords** antibody binding site definition and exposure • 2483: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This antibody was selected using a JRCSF fusion protein. Gorny et al. [2004] (antibody binding site definition and exposure)

No. 1172 **MAb ID** 2497 **HXB2 Location** Env Author Location Env (JR-CSF) **Epitope** Subtype B Neutralizing P Immunogen Species (Isotype) human

Ab Type gp120 V3

Research Contact Dr. Zolla-Pazner, Veterans Affairs Center, NY, NY. zollas01@endeavor.med.nyu.edu

References Gorny et al. 2004

Keywords antibody binding site definition and exposure • 2497: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This antibody was selected using a JRCSF fusion protein. Gorny et al. [2004] (antibody binding site definition and exposure)

No. 1173 **MAb ID** 2557 **HXB2 Location** Env Author Location Env (JR-CSF) **Epitope** Subtype A, CRF02\_AG Neutralizing P Immunogen HIV-1 infection Species (Isotype) human

Ab Type gp120 V3 Research Contact Dr. Zolla-Pazner, Veterans Affairs Center, NY,

NY. zollas01@endeavor.med.nyu.edu References Krachmarov et al. 2005; Gorny et al. 2004 Keywords antibody binding site definition and exposure, subtype comparisons, variant cross-

recognition or cross-neutralization

• 2557: Sera from subtype A infected individuals from Cameroon have antibodies that react strongly with subtype A and subtype B V3 loops in fusion proteins, and neutralize SF162 pseudotypes, while sera from 47 subtype B infected individuals reacted only with subtype B V3s. Sera from Cameroon did not neutralize primary A or B isolates, due to indirect masking by the V1/V2 domain rather than due to loss of the target epitope. 2557 was derived from a person infected with a clade

A or CRF02 virus, and binds to A and B V3 loops. Neutralization of JR-FL and SF162(UG V3) by anti-V3 MAbs 2557, 2558, 2601, but not subtype A primary isolates despite binding to the subtype A V3 loops, suggested masking by V1V2 blocking of neutralization by these antibodies. Krachmarov et al. [2005] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, subtype comparisons)

• 2557: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This antibody was selected Research Contact Dr. Zolla-Pazner, Veterans Affairs Center, NY, using a JRCSF fusion protein. Gorny et al. [2004]

No. 1174

**MAb ID** 2558

**HXB2 Location** Env

Author Location Env (92UG037)

**Epitope** 

Subtype A, CRF02\_AG

Neutralizing P

**Immunogen** 

Species (Isotype) human

Ab Type gp120 V3

Research Contact Dr. Zolla-Pazner, Veterans Affairs Center, NY,

NY. zollas01@endeavor.med.nyu.edu

References Krachmarov et al. 2005; Gorny et al. 2004

Country Uganda

Keywords antibody binding site definition and exposure, subtype comparisons, variant cross-

recognition or cross-neutralization

• 2558: Sera from subtype A infected individuals from Cameroon have antibodies that react strongly with subtype A and subtype B V3 loops in fusion proteins, and neutralize SF162 pseudotypes, while sera from 47 subtype B infected individuals reacted only with subtype B V3s. Sera from Cameroon did not neutralize primary A or B isolates, due to indirect masking by the V1/V2 domain rather than due to loss of the target epitope. 2557 was derived from a person infected with a clade A or CRF02 virus, and binds to A and B V3 loops. Neutralization of JR-FL and SF162(UG V3) by anti-V3 MAbs 2557, 2558, 2601, but not subtype A primary isolates despite binding to the subtype A V3 loops, suggested masking by V1V2 blocking of neutralization by these antibodies. Krachmarov et al. [2005] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, subtype comparisons)

• 2558: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This antibody was selected using an A clade fusion protein, 92UG037. It is unusual in that it is a V3 antibody selected for conformational aspects using an A clade virus, with a V3 GPGQ tip – clade B viruses are usually used and have GPGR tips. It cross-neutralizes and binds B

clade HIV SF162. Gorny et al. [2004] (antibody binding site definition and exposure)

**No.** 1175

**MAb ID** 2580

**HXB2 Location** Env

Author Location Env (JR-CSF)

**Epitope** 

Subtype B

Neutralizing P

**Immunogen** 

Species (Isotype) human

Ab Type gp120 V3

NY. zollas01@endeavor.med.nyu.edu

References Gorny et al. 2004

Keywords antibody binding site definition and exposure • 2580: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3

MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This antibody was selected using a JRCSF fusion protein. Gorny et al. [2004] (antibody

binding site definition and exposure)

**No.** 1176

MAb ID 391/95-D

**HXB2 Location** Env

Author Location Env

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

Research Contact S. Zolla-Pasner

References Guillon et al. 2002a

**Keywords** co-receptor, enhancing activity

391/95-D: This antibody was used to explore the sensitivity of chimeric envelope viruses to Ab-mediated enhancement or neutralization. V3 mediated enhancement and envelopes susceptible to enhancement used CCR5. Enhancement was CD4 dependent. Guillon et al. [2002a] (co-receptor, enhancing activity)

**No.** 1177

MAb ID 39F

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Subtype B

Neutralizing no

Immunogen

Species (Isotype)

Ab Type gp120 V3

Research Contact James Robinson, Tulane University, New Or-

leans, LA, USA

References Selvarajah et al. 2005; Pantophlet et al. 2004; Kwong et al. 2002; Grundner et al. 2002; Yang et al. 2002

**Keywords** antibody binding site definition and exposure, vaccine antigen design, vaccine-specific epitope characteristics

- 39F: Antigens were designed to attempt to target immune responses toward the IgG1b12 epitope, while minimizing antibody responses to less desirable epitopes. One construct had a series of substitutions near the CD4 binding site (GDMR), the other had 7 additional glycans (mCHO). The 2 constructs did not elicit b12-like neutralizing antibodies, but both antigens successfully dampened other responses that were intended to be dampened, while not obscuring b12 binding. V3 MAbs (447-52D, 19b, F245-B4e8 and 39F) bound to the GDMR antigen, but either did not bind or had diminished binding to mCHO. Selvarajah *et al.* [2005] (vaccine antigen design, vaccine-specific epitope characteristics)
- 39F: By adding N-linked glycosylation sites to gp120, epitope masking of non-neutralizing epitopes can be achieved leaving the IgG1b12 binding site intact. This concept was originally tested with the addition of four glycosylation sites, but binding to b12 was reduced. It was modified here to exclude the C1 N-terminal region, and to include only three additional glycosylation sites. This modified protein retains full b12 binding affinity and it masks other potentially competing epitopes, and does not bind to 21 other MAbs to 7 epitopes on gp120, including 39F. To inhibit 39F binding, Arg 304 and Lys 305 had to be changed to Ala. Pantophlet *et al.* [2004] (vaccine antigen design)
- 39F: HIV-1 gp160deltaCT (cytoplasmic tail-deleted) proteoliposomes (PLs) containing native, trimeric envelope glycoproteins from R5 strains YU2 and JRFL, and X4 strain HXBc2, were made in a physiologic membrane setting as candidate immunogens for HIV vaccines 2F5 bound to gp160deltaCT with a reconstituted membrane ten-fold better than the same protein on beads anti-CD4BS MAbs IgG1b12 and F105, A32 (C1-C4), C11 (C1-C5), and 39F (V3) MAbs bound gp160deltaCT PLs indistinguishably from gp160deltaCT expressed on the cell surface. Grundner et al. [2002]
- 39F: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-2999) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conforma-

tional or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong *et al.* [2002] (antibody binding site definition and exposure)

• 39F: Uncleaved soluble gp140 (YU2 strain, R5 primary isolate) can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif or using a T4 trimeric motif derived from T4 bacteriophage fibritin – stabilized oligomer gp140 delta683(-FT) showed strong preferential recognition by NAbs IgG1b12 and 2G12 relative to the gp120 monomer, in contrast to poorly neutralizing MAbs F105, F91, 17b, 48d, and 39F which showed reduced levels of binding, and C11, A32, and 30D which did not bind the stabilized oligomer. Yang *et al.* [2002]

**No.** 1178

**MAb ID** 4148d

**HXB2 Location** Env

**Author Location** Env

**Epitope** 

Subtype B

Neutralizing

Immunogen

Species (Isotype)

Ab Type gp120 V3

Research Contact Abraham Pinter, Public Health Research Institute, Newark, NJ, 07103. pinter@phri.org

References Pinter et al. 2004; Pinter et al. 1993b

**Keywords** antibody generation, variant cross-recognition or cross-neutralization

• 4148D: V1V2 was determined to be the region that conferred the neutralization phenotype differences between two R5-tropic primary HIV-1 isolates, JRFL and SF162. JRFL is resistant to neturalization by many sera and MAbs, while SF162 is sensitive. All MAbs tested, anti-V3, -V2, -CD4BS, and -CD4i, (except the broadly neutralizing MAbs IgG1b12, 2F5, and 2G12, which neutralized both strains), neutralized the SF162 pseudotype but not JRFL, and chimeras that exchanged the V1V2 loops transferred the neutralization phenotype. 5/6 anti-V3 MAbs, including 4148D, had similar binding affinity to soluble SF162 and JR-FL rgp120s, although the V3 loop differs at three positions (HigpgrafyTtgE for JR-FL and TigpgrafyAtgD for SF162). Pinter et al. [2004] (variant cross-recognition or cross-neutralization)

• 4148D: Pinter1993a first describes this MAb. Pinter *et al.* [1993b] (antibody generation)

**No.** 1179

MAb ID 55/68b

HXB2 Location Env

**Author Location** gp120 (300–315)

**Epitope** 

Neutralizing

Immunogen

Species (Isotype)

Ab Type gp120 V3

References Peet et al. 1998

 55/68b: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic – these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind, and anti-V3

MAb 55/68b binding was abrogated by V3 serine substitutions in the V3 loop – mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]

**No.** 1180 MAb ID 5G11 **HXB2 Location** Env Author Location gp120 **Epitope Neutralizing Immunogen** 

Species (Isotype) Ab Type gp120 V3

Research Contact S. Nigida and L. Arthur, NCI, Frederick, MD USA

References Moore & Sodroski 1996

• 5G11: Binds to conformation sensitive epitope in the V3 loop - reciprocal inhibition of other V3 loop MAbs - reciprocal enhancement of some C1-C5 MAbs (unusual for an anti-V3 MAb) and CD4 binding site MAbs - and enhances binding of V2 MAbs. Moore & Sodroski [1996]

No. 1181 **MAb ID** 6.1 **HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** Subtype B Neutralizing no Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

**Ab Type** gp120 V3

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

> References Gorny & Zolla-Pazner 2004; He et al. 2002 Kevwords review

- 6.1: This review provides summaries of Abs that bind to HIV-1 Research Contact Dr. Abraham Pinter, Public Health Research Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. 6.1 was non-neutralizing. Gorny & Zolla-Pazner [2004] (review)
- 6.1: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 – 3/4 V3 MAbs bound a similar linear epitope between positions 11-30 of the MN V3 loop (8E11/A8, 6.1, and 6.7), but only 8E11/A8 could weakly neutralize autologous strain SF162. He et al. [2002]

No. 1182 **MAb ID** 6.7 **HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** Subtype B Neutralizing no Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi adjuvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

**Ab Type** gp120 V3

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

> References Gorny & Zolla-Pazner 2004; He et al. 2002 Keywords antibody binding site definition and exposure, antibody generation, review

- 6.7: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. 6.7 was non-neutralizing. Gorny & Zolla-Pazner [2004] (review)
- 6.7: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 – 3/4 V3 MAbs bound a similar linear epitope between positions 11-30 of the MN V3 loop (8E11/A8, 6.1, and 6.7), but only 8E11/A8 could weakly neutralize autologous strain SF162. He et al. [2002] (antibody binding site definition and exposure, antibody generation)

**No.** 1183

**MAb ID** 8.27.3

**HXB2 Location** Env

Author Location gp120 (SF162)

**Epitope** Subtype B **Neutralizing** L Immunogen vaccine

> Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 V3

Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He et al. 2002 Keywords review, variant cross-recognition or crossneutralization

- 8.27.3: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, like 8.27.3; a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (variant crossrecognition or cross-neutralization, review)
- 8.27.3: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 - 1/4 V3 MAbs, 8.27.3, bound a discontinuous epitope that was broadly cross-reactive with B clade R5 and X4 strains (not E clade) and could neutralize autologous strain SF162. He et al. [2002]

No. 1184

**MAb ID** 8E11/A8

HXB2 Location Env

Author Location gp120 (SF162)

Epitope
Subtype B
Neutralizing L
Immunogen vaccine

Vector/Type: protein Strain: B clade SF162 HIV component: gp120 Adjuvant: Ribi ad-

juvant (MPL+TDM) (RIBI)

**Species (Isotype)** transgenic mouse (IgG2 $\kappa$ )

Ab Type gp120 V3

Research Contact Dr. Abraham Pinter, Public Health Research Institute, Newark, NJ, pinter@phri.org

References Gorny & Zolla-Pazner 2004; He *et al.* 2002 Keywords antibody binding site definition and exposure, antibody generation, autologous responses, review

- 8E11/A8: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 8E11/A8: Transgenic mice (strain XenoMouse G2) carrying human genes allowing production of fully human IgG2kappa MAbs were used to rapidly create a panel of anti-HIV gp120 MAb-producing hybridomas by immunization with HIV SF162 gp120 3/4 V3 MAbs bound a similar linear epitope between positions 11-30 of the MN V3 loop (8E11/A8, 6.1, and 6.7), but only 8E11/A8 could weakly neutralize autologous strain SF162. He *et al.* [2002] (antibody binding site definition and exposure, antibody generation, autologous responses)

**No.** 1185

**MAb ID** 9305

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L

Immunogen

Species (Isotype) mouse

**Ab Type** gp120 V3

Research Contact Du Pont, Wilmington DE

References McDougal et al. 1996

**No.** 1186

MAb ID A1g8

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

**Species (Isotype)** human ( $IgG1\lambda$ )

Ab Type gp120 V3

Research Contact James Robinson, Tulane University Med

School, New Orleans, LA, USA

**References** Cavacini *et al.* 2003; Cavacini *et al.* 2002

**Keywords** antibody interactions, co-receptor, variant cross-recognition or cross-neutralization

A1g8: The MAb B4e8 binds to the base of the V3 loop, neutralizes multiple primary isolates and was studied for interaction with other MAbs. B4e8 enhanced binding of CD4i MAbs 4.8d, 1.7b, and A1g8 to R5X4 virus 92HT593, but only of 48d to the R5 virus 92US660, and there was only a modest impact of the combination of B4e8 and CD4i MAbs on neutralization. Cavacini et al. [2003] (antibody interactions, co-receptor)

• A1g8: This study examined antibody interactions, binding and neutralization with a B clade R5 isolate (92US660) and R5X4 isolate (92HT593). Abs generally bound and neutralized the R5X4 isolate better than the R5 isolate. Anti-V3 MAb B4a1 increased binding of CD4i MAbs 48d, 17b and A1g8, but only A1g8 binding was increased by B4a1 to the R5 isolate. Additive affects on neutralization of the R5X4 isolate with B4a1 and CD4i MAbs was observed, presumably due to increased exposure of the CD4i binding site, but not for the R5 isolate. Antigp41 MAb F240 had a synergistic effect on neutralization with CD4i MAbs 48d and 17b, but not with A1g8 for the R5X4 virus. Cavacini *et al.* [2002] (antibody interactions, co-receptor, variant cross-recognition or cross-neutralization)

**No.** 1187

**MAb ID** AG1121 (1121)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen

Species (Isotype)

Ab Type gp120 V3

Research Contact AGMED, Inc, Bedford, MA, USA or ImmunoDiagnostics, Inc, Woburn, MA, USA

**References** Si et al. 2001; Cao et al. 1997b; Sullivan et al.

- AG1121: SHIV-HXBc2 is a neutralization sensitive non-pathogenic virus, and several *in vivo* passages through monkey's yielded highly pathogenic SHIV KU-1 HXBc2 and the KU-1 clone HXBc2P3.2 differ in 12 amino acids in gp160 substitutions in both gp120 and gp41 reduced the ability of sCD4, IgG1b12, F105 and AG1121 to Env achieve saturation and full occupancy, and neutralize KU-1 17b and 2F5 also bound less efficiently to HXBc2P3.2, although 2G12 was able to bind both comparably. Si *et al.* [2001]
- AG1121: Called 1121 Virus with the V1-V2 loop deleted was viable and more susceptible to neutralization by CD4i MAb 17b, and anti-V3 MAbs 1121, 9284, and 110.4, but not to and CD4BS MAb F105 or sCD4. Cao *et al.* [1997b]
- AG1121: Recognizes monomeric gp120 from T-cell adapted line HXBc2 and primary isolate 89.6 equally well, but 89.6 was three-fold less sensitive to neutralization by AG1121 than HXBc2. Sullivan et al. [1995]

**No.** 1188

MAb ID Ag1211

**HXB2 Location** Env

**Author Location** gp120 (V3) (JRFL)

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype)

**Ab Type** gp120 V3 **References** Kwong *et al.* 2002

Keywords antibody binding site definition and exposure

• Ag1211: Thermodynamics of binding to gp120 was measured using isothermal titration calorimetry for sCD4, 17b, b12, 48d, F105, 2G12 and C11 to intact YU2 and the HXBc2 core. The free energy of binding was similar, and not only CD4 but MAb ligands induced thermodynamic changes in gp120 that were independent of whether the core or the full gp120 protein was used. Non-neutralizing CD4BS and CD4i MAbs had large entropy contributions to free energy (mean: 26.1 kcal/mol) of binding to the gp120 monomer, except the potent CD4BS neutralizing MAb b6 had a much smaller value of 5.7 kcal/mol. High values suggest surface burial or protein folding and ordering of amino acids. Variable loop MAbs (L17, L78, 19b, 39F, Ag1211, D0142, and G3-2999) MAbs that bind to the N and C termini (211/c, A32, L100, P35, and C11) do not have dramatic entropy changes. These results suggest that while the trimeric Env complex has four surfaces, a non-neutralizing face (occluded on the oligomer), a variable face, a neutralizing face and a silent face (protected by carbohydrate masking), gp120 monomers further protect receptor binding sites by conformational or entropic masking, requiring a large energy handicap for Ab binding not faced by other anti-gp120 Abs. Kwong et al. [2002] (antibody binding site definition and exposure)

No. 1189
MAb ID B4a1
HXB2 Location Env
Author Location gp120 (V3)
Epitope
Subtype B
Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

Research Contact James Robinson, Tulane University Med School, New Orleans, LA, USA

> **References** Cavacini *et al.* 2003; Cavacini *et al.* 2002 **Keywords** antibody interactions, co-receptor, variant cross-recognition or cross-neutralization

- B4a1: The MAb B4e8 binds to the base of the V3 loop, neutralizes multiple primary isolates and was studied for interaction with other MAbs. The anti-V3 MAb B4a1 cross-competes with B4e8. Cavacini *et al.* [2003] (antibody interactions)
- B4a1: This study examined antibody interactions, binding and neutralization with a B clade R5 isolate (92US660) and R5X4 isolate (92HT593). Abs generally bound and neutralized the R5X4 isolate better than the R5 isolate. Anti-gp41 MAb F240 binding was not affected by the binding of the V3 loop MAb B4a1, but preincubation with F240 could enhance B4a1 binding of the R5 isolate. B4a1 reacts with many B clade isolates, and preincubation with sCD4 enhances binding to both the R5 and R5X4 isolates. B4a1 increased binding of CD4i MAbs 48d, 17b and A1g8, as well as CD4BS MAbs IgG1b12 and F105 to R5X4 virions, but only A1g8 and IgG1b12 binding was increased by B4a1 to the R5 isolate. Additive affects on neutralizaton of the R5X4 isolate with B4a1 and CD4i MAbs was observed, presumably due to increased exposure of the CD4i binding site, but not for the R5 isolate. B4a1 had an

additive affect on neutralization with 2G12 with the R5X4 virus but not the R5 virus, and did not impact 2F5 neutralization. Cavacini *et al.* [2002] (antibody interactions, co-receptor, variant cross-recognition or cross-neutralization)

**No.** 1190

**MAb ID** B4e8 (F425 B4e8)

**HXB2 Location** Env

Author Location gp120 (V3)

**Epitope** 

Subtype B

Neutralizing P

 $\begin{array}{c} \textbf{Immunogen} \ \ HIV\text{-}1 \ infection \\ \textbf{Species} \ (\textbf{Isotype}) \ \ \text{human} \ (\text{IgG2}\kappa) \end{array}$ 

Ab Type gp120 V3

Research Contact Lisa Cavacini, Beth Isreal Deconess Medical

Center, Boston MA, USA

References Lusso et al. 2005; Zwick et al. 2003; Liu et al.

2003; Cavacini et al. 2003

**Keywords** antibody binding site definition and exposure, antibody generation, antibody interactions, antibody sequence, variable domain, coreceptor, variant cross-recognition or cross-

neutralization

- B4e8: The epitope for the MAb D19 is conserved and embedded in V3. D19 is unique in that for R5 viruses, it was cryptic and did not bind without exposure to sCD4, and for X4 and R5X4 isolates it was constitutively exposed. It had an overlapping binding region with MAbs 447-52D, B4e8, and 268-D, but different reactivity patterns and fine specificity. While B4e8 and 447-52D could bind to the R5 virus BaL in the absence of sCD4, treatment with sCD4 did increase the binding of both B4e8 and 447-52D, but did not impact their ability to neutralize BaL. Lusso *et al.* [2005] (antibody binding site definition and exposure)
- B4e8: This MAb binds to the base of the V3 loop, and binds and neutralizes multiple primary isolates. The anti-V3 MAb B4a1 cross-competes with B4e8. B4e8 and 2G12 enhanced each others binding, and gave synergistic neutralization. B4e8 could neutralize R5X4 virus 92HT593 better than 2G12, while 2G12 was better at neutralizing R5 virus 92US660. B4e8 enhanced binding of CD4i MAbs 4.8d, 1.7b, and A1g8 to 92HT593, but only of 48d to the 92US660, and there was only a modest impact of the combination of B4e8 and CD4i MAbs on neutralization. CD4BS MAb IgG1b12 had no effect on B4e8 binding. Anti-gp41 MAb F240 inhibited B4e8 neutralization. Cavacini et al. [2003] (antibody binding site definition and exposure, antibody generation, antibody interactions, co-receptor, variant cross-recognition or cross-neutralization)
- B4e8: The effect of isotype (IgG1 and IgG3) and subtype (IgA) switching of parental F425B4e8 (IgG2) on HIV-1 binding and neutralization was investigated. IgG1-and IgA-F425B4e8 mutants showed virus-specific binding levels and TCLA SF2 isolate compared to the parental IgG2. Comparable levels of neutralization of primary isolates 92HT593 (R5X4) and 92US660 (R5) was achieved by all isotypes and subtypes of F425B4e8. Liu *et al.* [2003] (variant cross-recognition or cross-neutralization, antibody sequence, variable domain)

• B4e8: Called F425 B4e8. scFv 4KG5 reacts with a conformational epitope that is formed by the V1V2 and V3 loops and the bridging sheet (C4) region of gp120 and is influenced by carbohydrates. Of a panel of MAbs tested, only NAb b12 enhanced 4KG5 binding to gp120 JRFL. MAbs to the following regions diminished 4KG5 binding: V2 loop, V3 loop, V3-C4 region, CD4BS. MAbs directed against C1, CD4i, C5 regions didn't impact 4KG5 binding. These results suggest that the orientation or dynamics of the V1/V2 and V3 loops restricts CD4BS access on the envelope spike, and IgG1b12 can uniquely remain unaffected by these loops. This was one of the V3 MAbs used. Zwick et al. [2003] (antibody interactions)

No. 1191
MAb ID D27
HXB2 Location Env

**Author Location** gp120 (IIIB)

Epitope Neutralizing

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

**Species (Isotype)** mouse (IgG) **Ab Type** gp120 V3

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

**References** Sugiura *et al.* 1999; Otteken *et al.* 1996; Earl *et al.* 1994

- D27: A comparison of 25 gp120 specific, conformation dependent MAbs was done D27 is one of two MAbs labeled group Ca, that was type-specific for BH8 D27 fully blocked CD4 binding, and the deletion of the V3 loop abrogated binding. Sugiura *et al.* [1999]
- D27: Pulse label experiments of 4 MAbs (D20, D27, T20, and T22) binding to noncleavable gp160 revealed that these anti-CD4 MAbs bound with a delay, and that the epitope formed with a t 1/2 of about 10 minutes. Otteken *et al.* [1996]
- D27: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

**No.** 1192

MAb ID D47

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope Neutralizing** 

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB • HIV component: Env

Species (Isotype) mouse

Ab Type gp120 V3

Research Contact Patricia Earl, NIAID, NIH

References Salzwedel et al. 2000; Earl et al. 1997; Wyatt

et al. 1997; Otteken et al. 1996; Richardson

et al. 1996; Earl et al. 1994

**Keywords** antibody binding site definition and exposure, antibody generation, variant cross-recognition

or cross-neutralization

• D47: sCD4 can activate fusion between effector cells expressing Env and target cells expressing coreceptor (CCR5 or CXCR4) alone without CD4 – V3 MAb D47 is strain specific and can inhibit sCD4 mediated infection, but only of the closely related LAV Env, while anti-CD4i MAbs were broadly cross-neutralizing. Salzwedel *et al.* [2000] (variant cross-recognition or cross-neutralization)

- D47: Used for comparison in a study of gp41 antibodies D47 binds to a greater extent to cell surface expressed Env than any of 38 conformation dependent anti-gp41 MAbs. Earl *et al.* [1997]
- D47: Binds both gp120 and soluble gp120+gp41 complex efficiently, suggesting its gp120 epitope is not blocked by gp41 binding. Wyatt *et al.* [1997] (antibody binding site definition and exposure)
- D47: Pulse label experiments of MAb binding to noncleavable gp160 revealed that this anti-V3 MAb bound immediately and binding stayed constant through chase period. Otteken *et al.* [1996]
- D47: Used for capture of oligomeric Env for antigen capture ELISA binding of this antibody to oligomeric Env IIIB was not blocked by human sera from the US, consistent with a low prevalence of IIIB-like V3 strains. Richardson *et al.* [1996] (antibody binding site definition and exposure)
- D47: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994] (antibody generation)

No. 1193

MAb ID D56

**HXB2 Location** Env

**Author Location** gp120 (IIIB)

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade IIIB HIV component: oligomeric gp140

 $\textbf{Species (Isotype)} \ \ \text{mouse (IgG)}$ 

Ab Type gp120 V3

Research Contact P. Earl, National Institute of Allergy and Infectious Diseases, NIH, Bethesda, MD

References Sugiura et al. 1999; Earl et al. 1994

- D56: A comparison of 25 gp120 specific, conformation dependent MAbs was done D56 is one of two MAbs labeled group Ca, that was type-specific for BH8 D56 fully blocked CD4 binding, and the deletion of the V3 loop abrogated binding 12.5 ug/ml of D56 was required to achieve 50% neutralization of HIV-1 NL4-3. Sugiura et al. [1999]
- D56: Generated during a study of the influence of the oligomeric structure of Env in determining the repertoire of the Ab response. Earl et al. [1994]

No. 1194

MAb ID F5.5

HXB2 Location Env

Author Location gp120 (IIIB)

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype) mouse

Ab Type gp120 V3

Research Contact Hybridolabs, Institute Pasteur

References Altmeyer et al. 1999

• F5.5: A Semliki Forest virus (SFV) expression system carrying BX08 env was used to study the conformation of gp120 Env – intracytoplasmic gp120 was recognized by the anti-V3 MAbs K24 and F5.5, while gp120 at the plasma membrane was detected only by conformation dependent MAbs 2G12, 670-D and 694/98D and not V3 MAbs – expression in rat brain also showed that surface-expressed Env was recognized only by the conformation-dependent antibodies and not by anti-V3 antibodies. Altmeyer *et al.* [1999]

No. 1195

**MAb ID** G3-1472

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing

**Immunogen** 

Species (Isotype)

Ab Type gp120 V3

Research Contact M. Fung

References Moore & Sodroski 1996

G3-1472: Binds to carboxy-terminal side of the V3 loop – reciprocal binding inhibition with other anti-V3 and anti-C4 MAbs – reciprocal enhanced binding of some anti-V2 MAbs and anti-CD4 binding site MAbs – binding inhibited by anti-C4 MAbs. Moore & Sodroski [1996]

**No.** 1196

MAb ID K24

**HXB2 Location** Env

Author Location gp120 (IIIB)

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype) mouse

Ab Type gp120 V3

Research Contact Hybridolabs, Institute Pasteur

References Altmeyer et al. 1999

• K24: A Semliki Forest virus (SFV) expression system carrying BX08 env was used to study the conformation of gp120 env – intracytoplasmic gp120 was recognized by the anti-V3 MAbs K24 and F5.5, while gp120 at the plasma membrane was detected only by conformation dependent MAbs 2G12, 670-D and 694/98D and not V3 MAbs – expression in rat brain also showed that surface-expressed Env was recognized only by the conformation-dependent antibodies and not by anti-V3 antibodies. Altmeyer *et al.* [1999]

No. 1197

MAb ID TH1

**HXB2 Location** Env

Author Location gp120

**Epitope** 

**Neutralizing** L (MN, J)

Immunogen

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp120 V3

Research Contact Michael Fung, Tanox Biosystem, USA

References Gorny & Zolla-Pazner 2004; Yang et al. 1998;

D'Souza et al. 1995

**Keywords** assay development, review, variant crossrecognition or cross-neutralization

- TH1: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. TH1 neutralizes some TCLA strains. Gorny & Zolla-Pazner [2004] (review)
- TH1: A neutralization assay was developed based on heminested PCR amplification of the LTR (HNPCR) LTR-HNPCR consistently revealed HIV DNA and was shown to be a rapid, specific and reliable neutralization assay based on tests with 6 MAbs and 5 isolates. Yang et al. [1998] (assay development)
- TH1: Found to neutralize MN and JRCSF, but not two B subtype primary isolates, nor a D subtype primary isolate, by most labs in a multi-laboratory study involving 11 labs. D'Souza *et al.* [1995] (variant cross-recognition or cross-neutralization)

No. 1198

MAb ID anti-gp120/V3

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein, virus-like particle (VLP) Strain: A clade 94UG018 HIV component:

Gag, gp120, Nef, Pol

Species (Isotype) mouse (IgG)

**Ab Type** gp120 V3 **Research Contact** Intracel Co

References Buonaguro et al. 2001

Anti-V3: HIV-1 pr55 gag-based virus-like particles (VLP) carrying Nef and Pol open reading frames as well as gp120 of the clade A isolate 94UG018 were created using a Baculovirus expression system to package additional ORFS into the VLP – anti-V3 and anti-p24 antibodies were used to assess the expression levels and Gag and gp120-TM were found to be expressed at comparable levels on the VLP. Buonaguro *et al.* [2001]

**No.** 1199

MAb ID polyclonal

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: protein, virus-like particle (VLP)

Strain: B clade LAI HIV component:

CD4BS, Gag, V3

Species (Isotype) mouse

Ab Type gp120 V3

**References** Truong et al. 1996

Antibodies raised against recombinant anti-p55 virus-like particles with the p24 region 196-226 deleted, bearing inserts of either the V3 or the CD4BS regions of gp120 were studied – no neutralizing responses, weak Env, and strong Gag responses were elicited – the major homology region (MHR) and proximal sequences was found to be required for capsid assembly. Truong et al. [1996]

No. 1200
MAb ID polyclonal
HXB2 Location Env
Author Location gp120
Epitope
Neutralizing yes

Immunogen vaccine

Vector/Type: canarypox prime with recombinant protein boost Strain: B clade LAI, B clade MN, B clade SF2 HIV component: Gag, gp120, gp41, Pol Adjuvant: MF59

Species (Isotype) human

Ab Type gp120 V3

References Verrier et al. 2000

Serum Abs elicited by this vaccine reacted with V3 peptides from clades B, C, and F, reacted weakly with V3 peptides from clades A, D, G, and H, and did not react with V3 peptides from clades E and O – neutralizing activity against 5 of 14 primary isolates tested was observed, including one B clade X4 virus, two dualtropic B clade viruses (from clade B) and one clade B and one clade C R5 virus. Verrier et al. [2000]

No. 1201
MAb ID polyclonal
HXB2 Location Env
Author Location gp120 (303–325)
Epitope
Neutralizing no

References Sidorova 1999

Immunogen in vitro stimulation or selection

**Species (Isotype)** human (IgM) **Ab Type** gp120 V3

Polyspecific anti-MN-24 antibodies were raised through V3
peptide, MN-24 stimulation of human cells, followed by EBV
transformation: they react with homologous and heterologous
peptides and may be autoantibodies. Sidoroya [1999]

No. 1202
MAb ID polyclonal
HXB2 Location Env
Author Location
Epitope
Neutralizing
Immunogen
Species (Isotype) human

**Ab Type** gp120 V3

References Guevara et al. 2002

Viral RNA in serum and high titers of subtype C consensus V3 peptide binding Abs were the best independent predictors of mother to infant transmission of HIV-1 subtype C – NAb to subtype B HIV-1 (MN) was also correlated. Guevara *et al.* [2002]

No. 1203 MAb ID polyclonal HXB2 Location Env Author Location

> Epitope Subtype B Neutralizing L Immunogen vaccine

> > Vector/Type: HIV-1 captured on concavalin A-immobilized polystyrene nanospheres, Con A-NS Strain: B clade IIIB HIV component: gp120, heat-inactivated virus Adjuvant: concavalin A-immobilized polystyrene nanospheres

**Species (Isotype)** mouse (IgA) **Ab Type** gp120 V3

References Kawamura et al. 2002

 Vaginal fluids were collected after intravaginal immunization of BALB/c mice and analyzed for their anti-HIV-1 antibody levels using a IIIB-V3 ELISA and IIIB neutralization assay – HIV-1 specific IgG was undetectable but anti-HIV IgA antibody response was identified in the vaginal fluids of immunized mice with HIV concavalin A-immobilized polystyrene nanosheres. Kawamura et al. [2002]

MAb ID polyclonal
HXB2 Location Env

Author Location
Epitope
Subtype B
Neutralizing L
Immunogen vaccine

No. 1204

*Vector/Type:* peptide *Strain:* B clade 89.6P, B clade MN *HIV component:* Env *Adjuvant:* aluminum hydroxide, Cholera toxin (CT), GM-CSF, IL-12, IL-18, IL-1α

Species (Isotype) human (IgA, IgG1, IgG2a)

**Ab Type** gp120 V3 **References** Bradney *et al.* 2002

 The cytokine-adjuvant combination IL-1alpha, IL-12 and IL-18 were found to stimulate potent mucosal antibody responses upon intranasal immunization of mice – cholera toxin is the most widely used adjuvant, but is not safe for use in humans. Bradney et al. [2002]

No. 1205
MAb ID polyclonal
HXB2 Location Env
Author Location
Epitope
Subtype C
Neutralizing
Immunogen vaccine

Vector/Type: peptide Strain: multiple epitope immunogen HIV component: V3 Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (Isotype)** mouse **Ab Type** gp120 V3

References Hewer & Meyer 2002

• A synthetic peptide immunogen designated a multiple epitope immunogen (MEI) was generated by synthesizing peptides with mixtures of frequently found amino acids (>10%) from the C subtypes allowed in the synthetic peptide – when injected into mice, the C subtype MEI induced antibodies that recognized the immunogen and whole virus as an antigen in ELIZAs - sera from eight HIV positive South Africans recognized the MEI peptide in ELISA tests. Hewer & Meyer [2002]

**No.** 1206 MAb ID polyclonal **HXB2 Location** Env

Author Location gp120 (V3)

**Epitope** 

Subtype B, C, F

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Bongertz et al. 2003

Country Brazil

Keywords rate of progression, subtype comparisons

• Ab responses at diluations above 1:1000 against the consensus V3 loops of subtypes A, B, C, D, F, and Brazilian B and F, were detected in only 6/60 individuals infected with HIV by sexual exposure, while a significantly higher (38/46) reactivity and frequency of peptide recognition was observed in the plasma of IDUs. High Ab titers (> 1:10,000) were directed against V3B, V3Bbr and V3F peptides. The IDU group also displayed Research Contact J. Cordell, Institute for Cancer Research, Sutbroader NAb responses, in comparison to the sexually transmitted group. This may contribute to a slower disease progression in IDUs. Bongertz et al. [2003] (subtype comparisons, rate of progression)

No. 1207

MAb ID polyclonal

**HXB2 Location** Env

Author Location Env

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: adenovirus Strain: B clade HXB2/Bal HIV component: gp140ΔCFI, gp140ΔV1V2ΔCFImodifiedV3

Species (Isotype) guinea pig (IgG)

Ab Type gp120 V3

References Yang et al. 2004

Keywords co-receptor

• Neutralizing antibodies against V3 with greater breadth among B clade viruses were created in vaccinated guinea pigs using a combination gp140ΔV1V2 and shortened V3 loop envelope than using intact Envelope. The interior V3 glycosylation site was removed in the modification of V3. This change also caused the virus to become CXCR4 tropic. Yang et al. [2004] (co-receptor)

**No.** 1208

**MAb ID** 11/75a/21/41

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen

Species (Isotype)

Ab Type gp120 V3 discontinuous

References Peet et al. 1998; McKeating et al. 1992a

• 11/75a/21/41: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic – these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind, but anti-V3 MAb 11/75a/21/41 binding was dramatically diminished by V3 serine substitutions – mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al.

No. 1209

MAb ID 41.1 (ICR41.1i, ICR41. ICR 41.1i)

**HXB2 Location** Env

Author Location gp120 (HXB10)

**Epitope** 

Neutralizing L (HXB2)

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: gp120

Species (Isotype) rat (IgG2a)

Ab Type gp120 CD4i, gp120 V3 discontinuous

ton, Surrey, UK

References Heap et al. 2005a; Ugolini et al. 1997; Jeffs et al. 1996; Armstrong et al. 1996; Armstrong & Dimmock 1996; McLain & Dimmock 1994; Klasse et al. 1993a; McKeating et al. 1993b; McKeating et al. 1992a; Reitz et al. 1988

- 41.1: Called ICR 41.1i. Used as a positive control for enhanced MAb binding after sCD4 exposure – 41.1 binding to virions is increased 2-fold by sCD4. Heap et al. [2005a]
- 41.1: Viral binding inhibition by 41.1 was weakly correlated with neutralization (all other neutralizing MAbs tested showed some correlation except 2F5) Ugolini et al. [1997]
- 41.1: Called ICR41.1i IgG2c? Neutralization was affected if the Ab was added after the virus bound to the host cells at 24 degrees C or below. Armstrong & Dimmock [1996]
- 41.1: Called ICR41.1i Neutralization occurs by blocking a post-fusion internalization event, in contrast to MAb F58. Armstrong et al. [1996]
- 41.1: Deletion of the V1V2 regions did not affect anti-V3 Abs ability to bind when compared to intact rec gp120. Jeffs et al.
- 41.1: Called ICR41.1i Kinetics of neutralization studied no lag for 39.3b, while ICR 39.13g and ICR 41.1i have lags of 5 and 15 min respectively – neutralization mediated by 3 molecules of IgG per virion - most efficient at neutralization of the three MAbs studied - acts with multi-hit kinetics. McLain & Dimmock [1994]

tional changes in gp120 that confer neutralization resistance to conformationally sensitive neutralizing MAbs – neutralization efficiency of 41.1 is not affected. Klasse et al. [1993a]; Reitz et al. [1988]

**No.** 1210 MAb ID 55/45a/11 **HXB2 Location** Env Author Location gp120 **Epitope** Neutralizing Immunogen Species (Isotype)

Ab Type gp120 V3 discontinuous

References Peet et al. 1998

• 55/45a/11: The most variable amino acids in the V3 loop were replaced with serines to make the immunodominant V3 loop less immunogenic - these changes did not affect the ability of sCD4 or MAbs to V1/V2, C1 and C4 to bind, and anti-V3 MAb 55/45a/11 binding was only marginally diminished by V3 serine substitutions – mice injected with serine substituted gp120 had a reduced response relative to wildtype, and no enhanced immunogenicity of conserved regions. Peet et al. [1998]

No. 1211 **MAb ID** 1108 **HXB2 Location** Env Author Location Env (987) **Epitope** Subtype B Neutralizing P

Immunogen HIV-1 infection **Species (Isotype)** human (IgG1 $\lambda$ ) Ab Type gp120 V3 mimotope

References Gorny et al. 2004; Gorny & Zolla-Pazner 2004; Zolla-Pazner et al. 1999b; Zolla-Pazner et al. 1999a

**Keywords** antibody binding site definition and exposure, antibody generation, mimotopes, review

- 1108: This review provides summaries of Abs that bind to HIV-1 Env. There are many V3 MAbs, many neutralize some TCLA strains, and a subset can also neutralize some primary isolates. Gorny & Zolla-Pazner [2004] (review)
- 1108: V3 MAb neutralization is influenced by retaining the epitope, exposure on the intact virion, mobility during CD4induced conformational change, and affinity. Anti-V3 MAbs selected using V3 peptides neutralize less effectively than V3 MAbs selected using fusion proteins or gp120, suggesting antigenic conformation is important. This MAb was selected using V3 peptides. Gorny et al. [2004] (antibody binding site definition and exposure)
- 1108: Selected with peptide 987, a mimotope of anti-V3 MAb 447-D – MAb peptide reactivity pattern clustered with immunological related MAbs: 1108, 386, 268, 311, 257, 694.8 – the amino acids HI tended to be critical for reactivity in this group. Zolla-Pazner et al. [1999b] (antibody binding site definition and exposure, antibody generation, mimotopes)

• 41.1: The gp41 mutation 582(Ala to Thr) results in conformais ADGAWRSVHLGPGRGSGSGMGK. Zolla-Pazner et al. [1999a] (antibody binding site definition and exposure, antibody generation)

> **No.** 1212 MAb ID polyclonal **HXB2 Location** Env Author Location gp120 (IIIB)

> > **Epitope Neutralizing**

Immunogen vaccine

Vector/Type: peptide Strain: B clade MN HIV component: gp120 Adjuvant: Cholera toxin (CT)

Species (Isotype) rabbit

**Ab Type** gp120 V3-C4

References Zinckgraf et al. 1999

· Nasal mucosal immunization and boosting of HIV peptide and was superior for inducing serum IgG and vaginal secretory IgA compared to nasal immunization and vaginal boosting vaginal immunization and boosting resulted low serum IgG and vaginal IgA and a high vaginal IgG response. Zinckgraf et al. [1999]

No. 1213 MAb ID polyclonal **HXB2 Location** Env **Author Location Epitope** Neutralizing

Immunogen HIV-1 infection Species (Isotype) human (IgA, IgG) **Ab Type** gp120 V3, gp120 V4 References Skott et al. 1999

• IgA and IgG from 45 HIV+ individuals was studied – people with low CD4+ cell counts had decreased levels IgA in saliva sera and saliva IgA was primarily directed toward Env – peptide ELISA studies indicated that teh dominant IgA epitopes were the V4 region (aa 385-409) and the C-term part of the V3 loop (aa 325-344), while the IgG response was directed towards the tip of the loop (aa 308-325). Skott et al. [1999]

**No.** 1214 MAb ID polyclonal **HXB2 Location** Env Author Location gp41 **Epitope** 

> Subtype B Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: gp41

Species (Isotype) rabbit (IgG)

Ab Type gp41 alpha-helical hairpin intermediate

References Louis et al. 2003 Keywords vaccine antigen design

• Polyclonal Abs raised against soluble trivalently linked N35CCG-N13 and N34CCG, the internal trimeric core of the coiled-coil ectodomain, inhibit HIV-1 Env-mediated cell fusion at levels comparable to 2G12. Louis et al. [2003] (vaccine antigen design)

**No.** 1215

MAb ID 2G12 (c2G12)

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing LP

Immunogen

Species (Isotype)

Research Contact Herman Katinger, Inst. Appl. Microbiol. or Polymun Scientific Inc., Vienna, Austria,

References

No. 1216

**MAb ID** 1367

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

**Species (Isotype)** human ( $IgG1\lambda$ )

Ab Type gp41 cluster I

Research Contact Susan Zolla-Pazner

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Center)

References Gorny & Zolla-Pazner 2004; Nyambi et al. 2000; Gorny et al. 2000; Gorny & Zolla-

Pazner 2000; Nyambi et al. 1998

**Keywords** antibody binding site definition and exposure,

review, subtype comparisons

- 1367: One of 24 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster I region (aa 579 - 604). Only one of these has any neutralizing potential, clone 3. Gorny & Zolla-Pazner [2004] (review)
- 1367: A cluster I epitope that binds to rgp41 567-647, recognizes a peptide N51-C43 complex trimer of heterodimers that approximates the core of the fusogenic form of gp41 – this MAb doesn't react with either of the peptides N51 or C43 individually – MAbs 50-69 and 1367 had similar properties. Gorny & Zolla-Pazner [2000] (antibody binding site definition and exposure)
- 1367: Binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared - no MAb was oligomer specific, but gp41 MAb 50-69 bound with a 5 fold preference for the oligomer, while other gp41 MAbs (1367, 98-6, 167-D, 1281, 1342, and 1379) did not show a preference. Gorny et al. [2000] (antibody binding site definition and exposure)
- 1367: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 5 cluster I anti-gp41 MAbs which showed good cross clade reactivity – 1367 weakly bound to the majority of isolates - no neutralizing activity was observed when tested with 5 isolates, but 1367 did not bind well to these isolates. Nyambi et al. [2000] (subtype comparisons)

• 1367: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H - anti-gp41 Abs 98-6, 1367 and 1342 were not able to bind detectably with any of the viruses from any clade. Nyambi et al. [1998] (subtype comparisons)

No. 1217

MAb ID 7B2

HXB2 Location Env

Author Location gp41

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype)

Ab Type gp41 cluster I

References Haynes et al. 2005; Binley et al. 2003; Binley

et al. 1999

**Keywords** antibody binding site definition and exposure, vaccine antigen design

- 7B2: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 7B2 has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)
- 7B2: The SOS mutant envelope protein introduces a covalent disulfide bond between gp120 surface and gp41 transmembrane proteins into the R5 isolate JR-FL by adding cysteines at residues 501 and 605. Pseudovirions bearing this protein bind to CD4 and co-receptor bearing cells, but do not fuse until treatment with a reducing agent, and are arrested prior to fusion after CD4 and co-receptor engagement. gp41 Abs 7B2 and 2.2B did not neutralize in any format, WT, SOS, nor when added postbinding. Binley et al. [2003] (vaccine antigen design)
- 7B2: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits - a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen – SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 – SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 - nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120gp41 complexes. Binley et al. [1999] (antibody binding site definition and exposure)

No. 1218

MAb ID 126-6 (SZ-126.6)

HXB2 Location Env

Author Location gp41 (HXB2)

**Epitope** Subtype B Neutralizing no

Immunogen HIV-1 infection **Species (Isotype)** human ( $IgG2\kappa$ )

**Ab Type** gp41 cluster II

Research Contact Susan Zolla-Pazner (Zol-Med

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Center, NY, NY

References Gorny & Zolla-Pazner 2004; Finnegan et al. 2002; Nyambi et al. 2000; Gorny & Zolla-Pazner 2000; Hioe et al. 1997b; Earl et al. 1997; Binley et al. 1996; Chen et al. 1995; Eddleston et al. 1993; Xu et al. 1991; Robinson et al. 1991; Robinson et al. 1990b

Keywords antibody binding site definition and exposure, enhancing activity, kinetics, review, subtype comparisons, variant cross-recognition or cross-neutralization

- 126-6: NIH AIDS Research and Reference Reagent Program:
- 126-6: One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004]
- 126-6: Anti-gp41 MAbs were tested in a cell-cell fusion system to investigate the antigenic changes in gp41 during binding and fusion. Cluster I MAbs 50-69, F240, 240-D,3D6, and 246-D recognize a nonhelical hydrophobic region, positions 598-604, that forms a disulfide loop in the six-helix bundle. Cluster II MAbs 98-6 and 126-6 recognized residues 644-663 of gp41, a portion of the second heptad repeat. These MAbs were found to behave similarly, so 50-69 and 98-6 were used as representatives. Exposure of cluster I and cluster II epitopes required CD4 expression on HIV HXB2 Env expressing HeLa target cells, but not the CXCR4 co-receptor. Binding to CD4 exposed hidden cluster I and II epitopes. The MAbs were found to bind to gp120/gp41 complexes, not to gp41 after shedding of gp120, and were localized to at fusing-cell interfaces. Kinetic and binding results indicate that these MAbs are exposed in transitional structures during the fusion process, possibly the prehairpin intermediate prior to co-receptor binding, although other intermediate structures may be involved. They do not bind once syncytia begin to show extensive cyotplasmic mixing. These MAbs failed to inhibit fusion. The NAb 2F5 has a very different behavior in this study. Finnegan et al. [2002] (antibody binding site definition and exposure, kinetics)
- 126-6: This cluster II MAb binds to a conformational epitope in the region 644-663 – like most cluster II MAbs (126-6, 167-D, 1281, 1342, and 1379 all reacted similarly) it binds to a peptide N51-C43 complex trimer of heterodimers that approximates the core of the fusogenic form of gp41, but not to C43 nor to N51 alone – MAb 126-6 was biotinylated and used as a probe to determine that anti-gp41 MAb 50-69 bound the fusogenic form of the protein in liquid phase. Gorny & Zolla-Pazner [2000] (antibody binding site definition and exposure)
- 126-6: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 cluster II anti-gp41 MAbs

- of these 2F5, 167-D, 126-6, and 1281 bound across clades, but usually weakly, while 98-6 and 1342 had poor cross reactivity - Clade D isolates bound most consistently to cluster II MAbs. Nyambi et al. [2000] (variant cross-recognition or cross-neutralization, subtype comparisons)
- 126-6: Discontinuous epitope recognizing residues between 649-668 – designated cluster II – Fabs D5, D11, G1, T3, M12, M15, S6, S8, S9, S10 block binding. Binley et al. [1996] (antibody binding site definition and exposure)
- 126-6: One of several anti-gp41 MAbs that bind to a gp41maltose binding fusion protein designed to study the leucine zipper domain of gp41, showing that the construct has retained aspects of normal gp41 conformation. Chen et al. [1995] (antibody binding site definition and exposure)
- 126-6: Called SZ-126.6. Eddleston et al. [1993]
- 126-6: No enhancing or neutralizing activity. Robinson et al. [1991] (enhancing activity)
- 126-6: Specific for a conformational epitope. Xu et al. [1991] (antibody binding site definition and exposure)
- 126-6: No enhancing activity for HIV-1 IIIB. Robinson et al. [1990b] (enhancing activity)

No. 1219

**MAb ID** 1342

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

Ab Type gp41 cluster II

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

References Gorny & Zolla-Pazner 2004; Nyambi et al.

2000; Gorny et al. 2000; Gorny & Zolla-

Pazner 2000; Nyambi et al. 1998

Keywords antibody binding site definition and exposure, review, subtype comparisons

- 1342: One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- 1342: This cluster II MAb is a conformational epitope that binds in the region 644-663 – like most cluster II MAbs (126-6, 167-D, 1281, 1342, and 1379 all reacted similarly) it binds to a peptide N51-C43 complex trimer of heterodimers that approximates the core of the fusogenic form of gp41, but not to C43 nor to N51 alone. Gorny & Zolla-Pazner [2000] (antibody binding site definition and exposure)
- 1342: Binds within the region gp41 647-682 binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared - no MAb was oligomer specific, but gp41 MAb 50-69 bound with a 5 fold preference for the oligomer, while other gp41 MAbs (1367, 98-6, 167-D, 1281, 1342, and 1379) did not show a preference. Gorny et al. [2000] (antibody binding site definition and exposure)

- 1342: 26 HIV-1 group M isolates (clades A to H) were tested for binding to 47 MAbs, including 6 cluster II anti-gp41 MAbs of these 2F5, 167-D, 126-6, and 1281 bound across clades, but usually weakly, while 98-6 and 1342 had poor cross reactivity Clade D isolates bound most consistently to cluster II MAbs no neutralizing activity was observed when tested with 5 isolates, but 1342 did not bind to these isolates. Nyambi *et al.* [2000] (subtype comparisons)
- 1342: Using a whole virion-ELISA method, 18 human MAbs were tested for their ability to bind to a panel of 9 viruses from clades A, B, D, F, G, and H anti-gp41 Abs 98-6, 1367 and 1342 were not able to bind detectably with any of the viruses from any clade. Nyambi *et al.* [1998] (**subtype comparisons**)

No. 1220

**MAb ID** 1379

**HXB2 Location** Env

**Author Location** gp41

Epitope

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG1 $\lambda$ ) Ab Type gp41 cluster II

Research Contact Susan

t Susan Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu) (NYU Med.

Center)

References Gorny & Zolla-Pazner 2004; Gorny et al.

2000; Gorny & Zolla-Pazner 2000

**Keywords** antibody binding site definition and exposure, review

- 1379: One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- 1379: This cluster II MAb binds to a conformational epitope in the region 644-663 like most cluster II MAbs (126-6, 167-D, 1281, 1342, and 1379 all reacted similarly) it binds to a peptide N51-C43 complex trimer of heterodimers that approximates the core of the fusogenic form of gp41, but not to C43 nor to N51 alone. Gorny & Zolla-Pazner [2000] (antibody binding site definition and exposure)
- 1379: Binds within the region gp41 647-682 binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, but gp41 MAb 50-69 bound with a 5 fold preference for the oligomer, while other gp41 MAbs (1367, 98-6, 167-D, 1281, 1342, and 1379) did not show a preference. Gorny *et al.* [2000] (antibody binding site definition and exposure)

**No.** 1221

MAb ID 2.2B

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing no

**Immunogen** 

Species (Isotype)

Ab Type gp41 cluster II

Research Contact James Robinson, Tulane University, Tulane,

**References** Haynes *et al.* 2005; Binley *et al.* 2003; Schulke *et al.* 2002; Binley *et al.* 1999

**Keywords** antibody binding site definition and exposure, vaccine antigen design

- 2.2B: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. Haynes *et al.* [2005] (antibody binding site definition and exposure)
- 2.2B: The SOS mutant envelope protein introduces a covalent disulfide bond between gp120 surface and gp41 transmembrane proteins into the R5 isolate JR-FL by adding cysteines at residues 501 and 605. Pseudovirions bearing this protein bind to CD4 and co-receptor bearing cells, but do not fuse until treatment with a reducing agent, and are arrested prior to fusion after CD4 and co-receptor engagement. gp41 Abs 7B2 and 2.2B did not neutralize in any format, WT, SOS, nor when added postbinding. Binley *et al.* [2003] (vaccine antigen design)
- 2.2B: Ab binding characteristics of SOS gp140 were tested using SPR and RIPA SOS gp140 is gp120-gp41 bound by a disulfide bond NAbs 2G12, 2F5, IgG1b12, CD4 inducible 17b, and 19b bound to SOS gp140 better than uncleaved gp140 (gp140unc) and gp120 non-neutralizing MAbs 2.2B (binds to gp41 in gp140unc) and 23A (binds gp120) did not bind SOS gp140. Schulke *et al.* [2002] (vaccine antigen design)
- 2.2B: The MAbs with the broadest neutralizing activity, IgG1b12, 2G12 and 2F5, all have high affinity for the native trimer, indicating that they were raised in an immune response to the oligomer on the virion surface rather than dissociated subunits - a disulfide linked gp120-gp41 (SOS gp140) was created to mimic the native conformation of Env and explore its potential as an immunogen – SOS gp140 is recognized by NAbs IgG1b12, 2G12, and CD4-IgG2, and also by anti-V3 MAbs 19b and 83.1 - SOSgp140 is not recognized by C4 region MAbs that neutralize only TCLA strains, G3-42 and G3-519 - nor did it bind C11, 23A, and M90, MAbs that bind to gp120 C1 and C5, where it interacts with gp41 – MAbs that bind CD4 inducible epitopes, 17b and A32 were very strongly induced by CD4 in SOS gp140 - anti-gp41 MAbs that bind in the region that interacts with gp120, 7B2, 2.2B, T4, T15G1 and 4D4, did not bind to SOSgp140, in contrast to 2F5, which binds to the only gp41 epitope that is well exposed in native gp120-gp41 complexes. Binley et al. [1999] (vaccine antigen design)

**No.** 1222

MAb ID Fab D11 (D11)

HXB2 Location Env

**Author Location** gp41 (LAI)

Epitope

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

**Ab Type** gp41 cluster II

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, **Author Location** gp41 (LAI) antibody sequence, variable domain, review

- Fab D11: Called D11. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab D11: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody sequence, variable domain)

**No.** 1223

MAb ID Fab D5 (D5)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection **Species (Isotype)** human (IgG1 $\kappa$ )

**Ab Type** gp41 cluster II

References Gorny & Zolla-Pazner 2004; Binley et al. 1996

Keywords antibody binding site definition and exposure, antibody sequence, variable domain, review

- Fab D5: Called D5. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab D5: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody sequence, variable domain)

**No.** 1224

MAb ID Fab G1

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

**Ab Type** gp41 cluster II

References Gorny & Zolla-Pazner 2004; Binley et al. 1996

**Keywords** antibody binding site definition and exposure, antibody sequence, variable domain, review

- Fab G1: Called G1. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab G1: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody sequence, variable domain)

No. 1225

MAb ID Fab M10

**HXB2 Location** Env

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

**Ab Type** gp41 cluster II

References Parren et al. 1997b; Binley et al. 1996

- Fab M10: Does not bind to MN native oligomer, but does bind to both LAI and MN rgp120 and rgp140. Parren et al. [1997b]
- Fab M10: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 – conformation sensitive – variable regions sequenced. Binley et al. [1996]

**No.** 1226

**MAb ID** Fab M12 (M12)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp41 cluster II

References Gorny & Zolla-Pazner 2004; Binley et al. 1996

Keywords antibody binding site definition and exposure, antibody sequence, variable domain, review

- Fab M12: Called M12. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab M12: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody sequence, variable domain)

No. 1227

MAb ID Fab M15

**HXB2 Location** Env

**Author Location** gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

**Ab Type** gp41 cluster II

References Gorny & Zolla-Pazner 2004; Binley et al. 1996

Keywords review

- Fab M15: Called M15. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab M15: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 - conformation sensitive - variable regions sequenced. Binley et al. [1996]

**No.** 1228

**MAb ID** Fab S10 (S10)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**Ab Type** gp41 cluster II

References Gorny & Zolla-Pazner 2004; Binley et al.

1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, vari-

able domain, review

- Fab S10: Called S10. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab S10: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 – conformation sensitive – variable regions sequenced. Binley *et al.* [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

No. 1229

MAb ID Fab S6 (S6)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp41 cluster II

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain, review

- Fab S6: Called S6. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab S6: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 conformation sensitive variable regions sequenced. Binley *et al.* [1996] (antibody binding site definition and exposure, antibody interactions, antibody sequence, variable domain)

**No.** 1230

MAb ID Fab S8 (S8)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp41 cluster II

References Gorny & Zolla-Pazner 2004; Binley et al.

**Keywords** antibody binding site definition and exposure, antibody generation, antibody interactions, antibody sequence, variable domain, review

- Fab S8: Called S8. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (**review**)
- Fab S8: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody interactions, antibody sequence, variable domain)

**No.** 1231

MAb ID Fab S9 (S9)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp41 cluster II

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody interactions, antibody sequence, variable domain, review

- Fab S9: Called S9. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab S9: Binds to cluster II region competes with MAbs 126-6, Md-1 and D50 conformation sensitive variable regions sequenced. Binley *et al.* [1996] (antibody binding site definition and exposure, antibody generation, antibody interactions, antibody sequence, variable domain)

**No.** 1232

MAb ID Fab T3 (T3)

**HXB2 Location** Env

**Author Location** gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp41 cluster II

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

• Fab T3: Called T3. One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)

 Fab T3: Binds to cluster II region – competes with MAbs 126-6, Md-1 and D50 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

No. 1233

**MAb ID** Md-1 (MD-1)

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing no

Immunogen

**Species (Isotype)** human (IgG1 $\lambda$ )

**Ab Type** gp41 cluster II

Research Contact R. A. Myers State of Maryland Dept. of Health

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996; Chen *et al.* 1995; Myers *et al.* 1993

**Keywords** antibody binding site definition and exposure, review

- Md-1: NIH AIDS Research and Reference Reagent Program: 1223.
- Md-1: One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Md-1: Discontinuous epitope recognizing residues between 563-672, does not recognize cluster I disulfide bridge region reacts almost exclusively with trimers and tetramers on WB designated cluster II Fabs D5, D11, G1, T3, M12, M15, S6, S8, S9, S10 block binding. Binley *et al.* [1996] (antibody binding site definition and exposure)
- Md-1: Called MD-1 one of several anti-gp41 MAbs that bind to a gp41-maltose binding fusion protein designed to study the leucine zipper domain of gp41, showing that the construct has retained aspects of normal gp41 conformation. Chen *et al.* [1995] (antibody binding site definition and exposure)
- Md-1: Called MD-1 discontinuous epitope that binds in the N-terminal region – reacts exclusively with oligomer. Myers et al. [1993] (antibody binding site definition and exposure)

**No.** 1234

MAb ID Fab A9 (A9)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type gp41 cluster III

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, vari-

able domain, review

• Fab A9: Called A9. One of six Fabs in this database that bind to the gp41 cluster III region (a conformational epitope involving aa 619-648); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)

Fab A9: Binds to cluster III region – competes with MAb Md-1, but not MAbs 126-6 and D50 – conformation sensitive – variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

**No.** 1235

**MAb ID** Fab G15 (G15)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp41 cluster III

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

- Fab G15: Called G15. One of six Fabs in this database that bind to the gp41 cluster III region (a conformational epitope involving aa 619-648); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab G15: Binds to cluster III region competes with MAb Md-1, but not MAbs 126-6 and D50 conformation sensitive variable regions sequenced. Binley *et al.* [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

**No.** 1236

MAb ID Fab G5

**HXB2 Location** Env

**Author Location** gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**Ab Type** gp41 cluster III

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

- Fab G5: Called G5. One of six Fabs in this database that bind to the gp41 cluster III region (a conformational epitope involving aa 619-648); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab G5: Binds to cluster III region competes with MAb Md-1, but not MAbs 126-6 and D50 conformation sensitive variable regions sequenced. Binley *et al.* [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

**No.** 1237

MAb ID Fab L1 (L1)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

**Ab Type** gp41 cluster III

References Gorny & Zolla-Pazner 2004; Binley et al.

1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, vari-

able domain, review

- Fab L1: Called L1. One of six Fabs in this database that bind to the gp41 cluster III region (a conformational epitope involving aa 619-648); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab L1: Binds to cluster III region competes with MAb Md-1, but not MAbs 126-6 and D50 conformation sensitive variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

No. 1238

**MAb ID** Fab L11 (L11)

**HXB2 Location** Env

Author Location gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\kappa$ )

Ab Type gp41 cluster III

**References** Gorny & Zolla-Pazner 2004; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

- Fab L11: Called L11. One of six Fabs in this database that bind to the gp41 cluster III region (a conformational epitope involving aa 619-648); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab L11: Binds to cluster III region competes with MAb Md-1, but not MAbs 126-6 and D50 conformation sensitive variable regions sequenced. Binley et al. [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

**No.** 1239

MAb ID Fab L2 (L2)

**HXB2 Location** Env

**Author Location** gp41 (LAI)

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\kappa$ )

Ab Type gp41 cluster III

Research Contact P. Perrin and D. Burton (Scripps Research Institute, La Jolla, California

**References** Gorny & Zolla-Pazner 2004; Earl *et al.* 1997; Binley *et al.* 1996

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, review

- Fab L2: Called L2. One of six Fabs in this database that bind to the gp41 cluster III region (a conformational epitope involving aa 619-648); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- Fab L2: Binds to cluster III region competes with MAb Md-1, but not MAbs 126-6 and D50 conformation sensitive variable regions sequenced. Binley *et al.* [1996] (antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain)

**No.** 1240

**MAb ID** 1281 (1281-D)

**HXB2 Location** Env

**Author Location** gp41

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

**Species (Isotype)** human (IgG1 $\lambda$ )

**Ab Type** gp41 cluster II, gp41six-helix bundle

Research Contact Susan Zolla-Pazner (Zollas01@mcrcr6.med.nyu) (NYU Med.

Center)

**References** Gorny & Zolla-Pazner 2004; Follis *et al.* 2002; Golding *et al.* 2002b; Verrier *et al.* 

2001; Gorny et al. 2000; Gorny & Zolla-

Pazner 2000; Hioe et al. 1997b

**Keywords** antibody binding site definition and exposure, antibody interactions, review, variant cross-recognition or cross-neutralization

- 1281: One of 19 MAbs and Fabs in this database that bind to the highly immunogenic gp41 cluster II region (aa 644-663); none have neutralizing activity. Gorny & Zolla-Pazner [2004] (review)
- 1281: Alanine mutations were introduced into the N- and C-terminal α-helices of gp41 to destabilize interhelical packing interactions in order to study their inhibitory effect on viral infectivity. These mutations were shown to inhibit viral replication though affecting the conformational transition to the fusion-active form of gp41, and allow increased inhibition by gp41 peptides. 2F5 senstivity is increased in the mutated viruses, presumably because 2F5s neutralization activity is focused on the transition to the fusion active state. No other gp41 MAb against tested, including NC-1, 50-69D, 1281, 98-6D, 246-D and F240, neutralized the parental or the fusion-deficient mutated viruses. Follis et al. [2002] (antibody binding site definition and exposure)
- 1281: The fusion process was slowed by using a suboptimal temperature (31.5 C) to re-evaluate the potential of Abs targeting fusion intermediates to block HIV entry preincubation of E/T cells at 31.5 C enabled polyclonal anti-N-HR Ab and anti-six-helix bundle Abs to inhibit fusion, indicating six-helix bundles form prior to fusion 98-6 binds to a C-HR hairpin

epitope and blocks fusion when added to a 2 hour E/T preincubation at 31.5 C, but if added after 1 hour, doesn't inhibit – this is in contrast to six-helix bundle Abs 167-D and 1281 that inhibit more efficiently when added after one hour of incubation. Golding *et al.* [2002b] (antibody binding site definition and exposure)

- 1281: A panel of 12 MAbs was used to identify those that could neutralize the dual-tropic primary isolate HIV-1 89.6 six gave significant neutralization at 2 to 10 ug/ml: 2F5, 50-69, IgG1b12, 447-52D, 2G12, and 670-D six did not have neutralizing activity: 654-D, 4.8D, 450-D, 246-D, 98-6, and 1281 no synergy, only additive effects were seen for pairwise combinations of MAbs, and antagonism was noted between gp41 MAbs 50-69 and 98-6, as well as 98-6 and 2F5. Verrier et al. [2001] (antibody interactions)
- 1281: This cluster II MAb binds to a conformational epitope in the region 644-663 like most cluster II MAbs (126-6, 167-D, 1281, 1342, and 1379 all reacted similarly) it binds to a peptide N51-C43 complex trimer of heterodimers that approximates the core of the fusogenic form of gp41, but not to C43 nor to N51 alone. Gorny & Zolla-Pazner [2000] (antibody binding site definition and exposure)
- 1281: Binds within the region gp41 647-682 binding of panel of 21 MAbs to soluble oligomeric gp140 versus gp41 or gp120 monomers was compared no MAb was oligomer specific, but gp41 MAb 50-69 bound with a 5 fold preference for the oligomer, while other gp41 MAbs (1367, 98-6, 167-D, 1281, 1342, and 1379) did not show a preference. Gorny *et al.* [2000] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization)
- 1281: Called 1281-D: Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe *et al.* [1997b] (variant cross-recognition or cross-neutralization)

**No.** 1241

MAb ID Chessie 8

**HXB2 Location** Env

Author Location gp41

Epitope

Neutralizing

Immunogen

Species (Isotype) mouse (IgG)

Ab Type gp41 cytoplasmic domain

Research Contact G. Lewis

References Smith-Franklin *et al.* 2002; Rovinski *et al.* 

1995; Poumbourios et al. 1995; Lewis et al. 1991

that HIV-1 antibody and Fcgamma receptors can trap virus on the surface of follicular dendritic cells (FDC)'s and extend the period of infectivity – blocking the FDC-Fcgamma receptor killing the FDC cell reduced their ability to maintain infectivity, and FDC cells seemed to stabilize viral particles and decrease gp120 shedding. Smith-Franklin *et al.* [2002]

• Chessie 8: This Ab was used in an *in vitro* study demonstrating

 Chessie 8: Used to precipitate gp160 in immunoblots in a study examining the feasibility of using unprocessed gp160 glycoprotein as an immunogen. Rovinski et al. [1995]

**No.** 1242

**MAb ID** 8F102

**HXB2 Location** Env

Author Location gp120

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain: B clade HXB2 HIV component: gp120

Species (Isotype) mouse (IgG)

Ab Type gp120-CD4 complex

References DeVico et al. 1995

8F102: MAbs specifically reactive to crosslinked gp120 and CD4 were derived (8F101, 8F102) – conformation dependent – competition studies indicate the epitope is immunogenic in infected humans. DeVico et al. [1995]

**No.** 1243

MAb ID CG-10 (CG10)

**HXB2 Location** Env

Author Location gp120 (IIIB)

Epitope

Subtype B

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex Strain:

B clade IIIB HIV component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120-CD4 complex

Ab Type gp120-CD4 complex

Research Contact Jonathan Gershoni, Tel Aviv University, Isreal

**References** Enshell-Seijffers *et al.* 2003; Finnegan *et al.* 2001; Oscherwitz *et al.* 1999a; Sullivan *et al.* 

1998b; Rizzuto *et al.* 1998; Lee *et al.* 1997; Wu *et al.* 1996; Gershoni *et al.* 1993

**Keywords** antibody binding site definition and exposure, computational epitope prediction, structure

• CG10: Using 17b MAb to select peptides from a combinatorial library, and analyzing the peptides using a novel discontinuous epitope reconstruction program, enabled epitope prediction. Segments of gp120 were reconstructed as an antigenic protein mimetic recognized by 17b. Comparisons then were made with a similar prediction of contact residues for CG10, a CD4i MAb that competes with 17b, but has a distinct binding site. Enshell-Seijffers *et al.* [2003] (antibody binding site definition and exposure, computational epitope prediction, structure)

- CG-10: Called CG10. Using a cell-fusion system, it was found CD4i antibodies 17b, 48d, and CG10 reacted faintly with Env expressing HeLA cells even in the absence of sCD4 or CD4 expressing target cells. Reactivity increased after sCD4 addition, but not after CD4 expressing target cell addition, and binding was not increased at the cell-to-cell CD4-Env interface. This suggests the CD4i co-receptor binding domain is largely blocked at the cell-fusion interface, and so CD4i antibodies would not be able access this site and neutralize cell-mediated viral entry. Finnegan *et al.* [2001] (antibody binding site definition and exposure)
- CG-10: Called CG10 disrupts gp120-CCR5 interaction and competes with MAb 17b –binds near the conserved bridging sheet of gp120 mutations in positions K/D 121, T/D 123, K/D 207, K/D 421, Q/L 422, Y/S 435, M/A 434, K/A 432 and I/S 423 result in a 70% reduction in CG10 binding. Rizzuto et al. [1998]
- CG-10: Called CG10 CD4BS MAb 15e competes with CG-10 binding, probably due to the disruption of CD4-gp120 by 15e CD4i MAbs 17b and 48d compete and the binding sites may overlap MAb A32 enhances binding of 17b, 48d and CG10 MAbs C11, 2G12 and 212A do not affect CG10 binding CG-10 can bind gp120 with V1/V2 and V3 deleted HXBc2 mutations Delta 119-205, 314 G/W, 432 K/A, 183,184 PI/SG decrease CG-10 recognition, HXBc2 mutations Delta 298-327 (V3), 384 Y/E, 298 R/G, 435 Y/S enhance recognition the CD4 contribution to the CG10 epitope maps to the CD4 CDR2-like loop CG10 can neutralize HIV-1 in the presence of sCD4 even though it does not do so in the context of cell surface CD4 binding to gp120. Sullivan et al. [1998b]
- CG-10: Called CG10 Promotes envelope mediated cell fusion between CD4+ cells and cells infected with either T-cell and macrophage tropic viruses – infection of HeLa CD4+ (MAGI) cells by HIV-1 LAI, ELI1, and ELI2 strains was increased two-to four-fold in the presence of CG10. Lee *et al.* [1997]
- CG-10: Called CG10 MIP-1alpha binding to CCR-5 expressing cells can be inhibited by gp120-sCD4, and MAb CG10 does not block this inhibition. Wu et al. [1996]
- CG-10: Reacts exclusively with sCD4-gp120 complex, not with sCD4 or gp120 alone. Gershoni *et al.* [1993]

**No.** 1244

MAb ID CG-25

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex HIV

component: gp120

**Species (Isotype)** mouse (IgG1)

**Ab Type** gp120-CD4 complex

**References** Gershoni *et al.* 1993

 CG-25: Reacts preferentially with sCD4-gp120, also with sCD4, not with gp120. Gershoni et al. [1993]

**No.** 1245

MAb ID CG-4 (CG4)

**HXB2 Location** Env **Author Location** gp120

Epitope

Neutralizing no

Immunogen vaccine

Vector/Type: sCD4-gp120 complex HIV

component: gp120

Species (Isotype) mouse (IgG1)

**Ab Type** gp120-CD4 complex

blocked at the cell-fusion interface, and so CD4i antibodies Research Contact Jonathan Gershoni, Tel Aviv University, Isreal would not be able access this site and neutralize cell-mediated References Gershoni *et al.* 1993

• CG-4: Reacts with gp120 and sCD4-gp120 complex, not with sCD4. Gershoni *et al.* [1993]

**No.** 1246

MAb ID CG-76

**HXB2 Location** Env

**Author Location** gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex HIV

component: gp120

Species (Isotype) mouse (IgG1)

Ab Type gp120-CD4 complex

**References** Gershoni *et al.* 1993

 CG-76: Reacts equally well with sCD4-gp120 and sCD4, but not with purified gp120. Gershoni et al. [1993]

**No.** 1247

MAb ID CG-9

HXB2 Location Env

Author Location gp120

**Epitope** 

Neutralizing L

Immunogen vaccine

Vector/Type: sCD4-gp120 complex HIV

component: gp120

Species (Isotype) mouse (IgG1)

**Ab Type** gp120-CD4 complex

**References** Gershoni et al. 1993

 CG-9: Reacts preferentially with sCD4-gp120, also with sCD4, not with gp120. Gershoni et al. [1993]

**No.** 1248

**MAb ID** 105-518

HXB2 Location Env

Author Location gp41 (608–637 HAM112, O group)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: O group

HAM112 HIV component: gp160

**Species** (**Isotype**) mouse ( $IgG1\kappa$ )

Ab Type immunodominant region

References Scheffel et al. 1999

• 101-518: Overlapping peptides based on group O HAM112 Env were tested for MAb reactivity. Scheffel *et al.* [1999]

**No.** 1249

MAb ID 31A1

**HXB2 Location** Env

**Author Location** gp41 **Epitope** 

Neutralizing no

Immunogen in vitro stimulation or selection

**Species (Isotype)** human ( $IgM \kappa/\lambda$ )

Ab Type p24+gp41

References Pollock et al. 1989

 31A1: Denatured virus was used for *in vitro* stimulation to generate Abs – Reacts with both p24 and gp41. Pollock *et al*. [1989]

**No.** 1250

**MAb ID** 39A64

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing no

Immunogen in vitro stimulation or selection

**Species** (**Isotype**) human ( $\operatorname{IgM} \kappa/\lambda$ )

**Ab Type** p24+gp41

References Pollock et al. 1989

• 39A64: Denatured virus was used for *in vitro* stimulation to generate Abs – Reacts with both p24 and gp41. Pollock *et al.* [1989]

No. 1251

**MAb ID** 39B86

**HXB2 Location** Env

Author Location gp41

**Epitope** 

Neutralizing no

Immunogen in vitro stimulation or selection

**Species (Isotype)** human ( $IgM \kappa/\lambda$ )

**Ab Type** p24+gp41

References Pollock et al. 1989

 39B86: Denatured virus was used for *in vitro* stimulation to generate Abs – Reacts with both p24 and gp41. Pollock *et al*. [1989]

No. 1252

**MAb ID** 9303

HXB2 Location Env

**Author Location** gp41

**Epitope** 

Neutralizing no

Immunogen

Species (Isotype) mouse

Ab Type p24+gp41

Research Contact Du Pont

References McDougal et al. 1996

**No.** 1253

MAb ID NC-1

**HXB2 Location** Env

Author Location gp41 (IIIB)

Immunogen vaccine

Epitope

Neutralizing

Vector/Type: peptide Strain: B clade IIIB

HIV component: gp41

Species (Isotype) mouse (IgG2a)

**Ab Type** gp41six-helix bundle

Research Contact S. Jiang, New York Blood Center, NY, NY

References de Rosny et al. 2004a; de Rosny et al. 2004b;

Follis et al. 2002; Yang et al. 2002; Yang et al.

2000; Jiang et al. 1998

**Keywords** antibody binding site definition and exposure, antibody generation, antibody interactions, subtype comparisons, variant cross-

recognition or cross-neutralization

• NC-1: The MAb 2F5 binds to the C-heptad and is neutralizing, but the MAb D50 binds to the C-heptad and is not neutralizing. 2F5 binds preferentially to native gp41 prior to receptor activation. Trapped fusion-intermediates suggest 2F5 remains present shortly after gp120 triggering by CD4, but may be lost by the time the six-helix bundle is formed. 2F5 neutralization seems to block a later step of the fusion process, but it does not inhibit binding of NC-1, a MAb specific for the six-helix bundle, so it does not prevent formation of the six-helix bundle. The results are most consistent with 2F5 inhibiting a post-fusion-intermediate step. de Rosny *et al.* [2004b] (antibody binding site definition and exposure, antibody interactions)

- NC-1: The mechanism of 2F5 neutralization was explored, and experiments suggest it is due to interference with a late step in viral entry. 2F5 does not block six-helix bundle formation, as 2F5 prebinding does not inhibit NC-1 binding, a MAb that binds specifically to the six-helix bundle. de Rosny *et al.* [2004a] (antibody binding site definition and exposure)
- NC-1: Alanine mutations were introduced into the N- and C-terminal alpha-helices of gp41 to destabilize interhelical packing interactions in order to study their inhibitory effect on viral infectivity. These mutations were shown to inhibit viral replication though affecting the conformational transition to the fusion-active form of gp41, and allow increased inhibition by gp41 peptides. 2F5 senstivity is increased in the mutated viruses, presumably because 2F5s neutralization activity is focused on the transition to the fusion active state. No other gp41 MAb against tested, including NC-1, 50-69D, 1281, 98-6D, 246-D and F240, neutralized the parental or the fusion-deficient mutated viruses. Follis *et al.* [2002] (antibody binding site definition and exposure)
- NC-1: Uncleaved soluble gp140 can be stabilized in an oligomer by fusion with a C-term trimeric GCN4 motif (gp140delta683(-/GCN4)) or using a T4 trimeric motif derived from T4 bacteriophage fibritin (gp140delta683(-/FT)) NC-1 binds to 15% of the GCN4 motif trimers, but this was significantly reduced for the T4 fibritin stabilized structures, indicating little is in the six-helix bundle, fusogenic conformation. Yang *et al.* [2002] (antibody binding site definition and exposure)
- NC-1: A combination of gp41 fusion with the GNC4 trimeric sequences and disruption of the YU2 gp120-gp41 cleavage site resulted in stable gp140 trimers (gp140-GNC4) approximately 16% of the gp140(-GNC4) stabilized trimer recognized by pooled sera was precipitated by NC-1, indicating that at a fraction assumes a fusogenic gp41 six-helix bundle conformation gp140(-) monomers were not able to bind to the NC-1, nor was gp130(-/GCN4) glycoprotein, consistent with the expectation that the absence of C34 helices would preclude

formation of the six-helix bundle. Yang *et al.* [2000] (antibody binding site definition and exposure)

• NC-1: Ab elicited in response to immunization with N36(L6)C34, a peptide that folds into a six helix bundle like gp41 – NC-1 binds to the surface of HIV-1 infected cells only in the presence of sCD4, recognizing the fusogenic core structure – binding affinity was decreased by point mutations that disrupt core formation and abolish membrane fusion activity, (I573P and I573A) – NC-1 can recognize discontinuous epitopes from B clade isolate SC, but not E clade strain N243, O group strain GAB, or HIV-2 ROD. Jiang *et al.* [1998] (antibody binding site definition and exposure, antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons)

## IV-C-18 Nef Antibodies

No. 1254

MAb ID 4H4

HXB2 Location Nef (1–33)

**Author Location** Nef (1–33 IIIB)

Epitope MGGKWSKSSVVGWPTVRERMRRAPTVRERMRR-

AEPAADGVGAA

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade IIIB

HIV component: Nef

Species (Isotype) human (IgG1)

References Otake et al. 1994

• 4H4: This MAb, elicited by vaccination with a Nef fusion protein, could not detect Nef protein on the cell surface – C-term anti-Nef Abs could. Otake *et al.* [1994]

No. 1255

MAb ID polyclonal

HXB2 Location Nef (9–24)

**Author Location** Nef (9–24)

Epitope SVIGWLTVRERMRRAE

Neutralizing no

Immunogen vaccine

Vector/Type: DNA Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG)

References Tahtinen et al. 2001

• BALB/c mice were immunized with a pBN-vector expressing HIV-1 nef, rev, or tat genes – DNA loaded onto gold microparticles was delivered using a gene gun, and DNA dissolved in saline was given intradermally or intramuscularly – Nef gene gun immunized mice showed the strongest and most long-lasting (6 months) Ab, CTL and proliferative responses – the highest IgG1/IgG2a ratio was observed in the gene gun immunized mice – three Ab binding sites were found in Nef using peptide mapping, although some sera reacted only to complete Nef – Rev- or-Tat immunized mice did not generate an Ab response. Tahtinen *et al.* [2001]

**No.** 1256 **MAb ID** 13/042

**HXB2 Location** Nef (11–20)

**Author Location** Nef (11–24 BH10)

**Epitope** VGWPTVRERM

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse

References Schneider et al. 1991

 13/042: Epitope mapped by overlapping decapeptides – core: TVRERM. Schneider et al. [1991]

No. 1257

MAb ID 13/035

HXB2 Location Nef (15-24)

Author Location Nef (11-24 BH10)

Epitope TVRERMRRAE

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse

References Schneider et al. 1991

• 13/035: Epitope mapped by overlapping decapeptides – core: TVRERM. Schneider *et al.* [1991]

No. 1258

MAb ID A6

HXB2 Location Nef (18-26)

Author Location Nef (18-26 NL-432)

**Epitope** ERMRRAEPA?

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: Nef Adjuvant: Complete

Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgM)

References Otake et al. 1997

Keywords antibody binding site definition and exposure,

antibody generation

A6: BALB/c mice were immunized with Nef protein to create a series of anti-Nef antibodies. A6 bound to the peptide spanning amino acids 18-26; we inferred the amino acids from the positions in the NL-43 strain. A6 also bound to the complete Nef protein. Otake *et al.* [1997] (antibody binding site definition and exposure, antibody generation)

**No.** 1259

MAb ID AM5C6

HXB2 Location Nef (28–43)

**Author Location** Nef (28–43 BH10)

Epitope DGVGAASRDLEKHGAI+KAAVDLSHFLK

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse

References Maksiutov et al. 2002; Schneider et al. 1991

- AM5C6: This epitope is similar to a fragment of the human protein vascular endothelial growth factor C, AEPDAGEATAYASKDLE. Maksiutov *et al.* [2002]
- AM5C6: Epitope mapped by overlapping decapeptides core: SRDL also reacts with Nef(78-92). Schneider *et al.* [1991]

**No.** 1260

MAb ID AM5C6

HXB2 Location Nef (28-43)

**Author Location** Nef (28–43 BH10)

Epitope DGVGAASRDLEKHGAI+KAAVDLSHFLK

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse

References Maksiutov et al. 2002; Schneider et al. 1991

- AM5C6: This epitope is similar to a fragment of the human protein vascular endothelial growth factor C, AEPDAGEATAYASKDLE. Maksiutov et al. [2002]
- AM5C6: Epitope mapped by overlapping decapeptides core: KAAVDL - also reacts with Nef(28-43). Schneider et al.

No. 1261

MAb ID A7

HXB2 Location Nef (28-45)

Author Location Nef (28-45 NL-432)

Epitope DGVGAVSRDLEKHGAITS?

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgG1)

References Otake et al. 1997

antibody generation

• A7: BALB/c mice were immunized with Nef protein to create a series of anti-Nef antibodies. A7 bound to the peptide spanning amino acids 28-45; we inferred the amino acids from the positions in the NL-43 strain. A7 did not bind to the complete Nef protein. Otake et al. [1997] (antibody binding site definition and exposure, antibody generation)

**No.** 1262

MAb ID 25/03

HXB2 Location Nef (30-43)

**Author Location** Nef (30–43 BH10)

Epitope VGAASRDLEKHGAI

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse

References Maksiutov et al. 2002; Schneider et al. 1991

- 25/03: This epitope is similar to a fragment of the human protein vascular endothelial growth factor C, AEPDAGEATAYASKDLE. Maksiutov et al. [2002]
- 25/03: Epitope mapped by overlapping decapeptides core: ASRDLEK. Schneider et al. [1991]

No. 1263

MAb ID 26/76

HXB2 Location Nef (30-43)

**Author Location** Nef (30–43 BH10)

Epitope VGAASRDLEKHGAI

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse

References Maksiutov et al. 2002; Schneider et al. 1991

- 26/76: This epitope is similar to a fragment of the human protein vascular endothelial growth factor C, AEPDAGEATAYASKDLE. Maksiutov et al. [2002]
- 26/76: Epitope mapped by overlapping decapeptides core: SRDLEK. Schneider et al. [1991]

No. 1264

MAb ID 3F2

HXB2 Location Nef (31–40)

**Author Location** Nef (31–40 BRU)

Epitope GAASRDLEKH

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG1)

References Maksiutov et al. 2002; Ranki et al. 1995; Saito et al. 1994; Ovod et al. 1992

- 3F2: UK Medical Research Council AIDS reagent: EVA3067.1.
- 3F2: This epitope is similar to a fragment of the human protein vascular endothelial growth factor C, AEPDAGEATAYASK-DLE. Maksiutov et al. [2002]
- 3F2: Faintly cross-reactive with astrocytes of uninfected control samples. Ranki et al. [1995]
- Keywords antibody binding site definition and exposure, 3F2: Reacted with Nef from different HIV-1 strains (BRU, IIIB, RF, MN). Ovod et al. [1992]

**No.** 1265

MAb ID 3D12

HXB2 Location Nef (31–50)

Author Location Nef (31–50 BRU)

Epitope GAASRDLEKHGAITSSNTAA

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: Nef

**Species (Isotype)** mouse (IgG1)

References Maksiutov et al. 2002; Ranki et al. 1995; Saito et al. 1994; Ovod et al. 1992

- 3D12: There is an anti-RT MAb that also has this name (see.
- 3D12: UK Medical Research Council AIDS reagent: EVA3067.2.
- 3D12: This epitope is similar to a fragment of the human protein vascular endothelial growth factor C, AEPDAGEATAYASKDLE. Maksiutov et al. [2002]
- 3D12: One of four antibodies used in combination to show HIV Nef protein expressed in astrocytes from 7/14 brain samples from HIV+ individuals - Nef expression associated with dementia. Ranki et al. [1995]
- 3D12: Over-expression of Nef in astrocytes from postmortem pediatric CNS tissues. Saito et al. [1994]
- 3D12: Reacted with Nef from different HIV-1 strains (BRU, IIIB, RF, MN). Ovod et al. [1992]

**No.** 1266

MAb ID polyclonal

HXB2 Location Nef (33-65)

Author Location Nef (32-64 LAI, BRU)

Epitope ASRDLEKHGAITSSNTAATNAACAWLEAQEEEE

Subtype B

Neutralizing

Immunogen HIV-1 infection, vaccine

Vector/Type: protein, PLG microparticle Strain: B clade BRU, B clade LAI HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA), PLG

Species (Isotype) mouse (IgG1)

References Maksiutov et al. 2002; Moureau et al. 2002

- This epitope is similar to a fragment of the human protein vascular endothelial growth factor C, AEPDAGEATAYASKDLE. Maksiutov et al. [2002]
- Nef encapsulated in poly(DL-lactide-co-glycolide) (PLG) had a more prolonged Ab response than Nef in PBS or in Freund's adjuvant (CFA), still strong after 7 months the response was predominantly IgG1, a Th2 immune response three linear epitopes, Nef 32-64, 118-167, and 185-205, were frequently recognized by the sera of mice immunized with NefPLG or Nef-CFA, but not after immunization with Nef in PBS, which seemed to preferentially stimulate an Ab response to conformational epitopes. Moureau et al. [2002]

No. 1267

MAb ID polyclonal

HXB2 Location Nef (49–64)

Author Location Nef (49-64)

Epitope AATNAACAWLEAQEEE

Neutralizing no Immunogen vaccine

Vector/Type: DNA Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG)

References Tahtinen et al. 2001

• BALB/c mice were immunized with a pBN-vector expressing HIV-1 nef, rev, or tat genes – DNA loaded onto gold microparticles was delivered using a gene gun, and DNA dissolved in saline was given intradermally or intramuscularly – Nef gene gun immunized mice showed the strongest and most longlasting (6 months) Ab, CTL and proliferative responses – the highest IgG1/IgG2a ratio was observed in the gene gun immunized mice – three Ab binding sites were found in Nef using peptide mapping, although some sera reacted only to complete Nef – Rev- or-Tat immunized mice did not generate an Ab response. Tahtinen *et al.* [2001]

**No.** 1268

**MAb ID** 3G12

**HXB2 Location** Nef (51–71)

**Author Location** Nef (51–71 BRU)

Epitope TNAACAWLEAQEEEEVGFPVT

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG2a)

References Ovod et al. 1992

 3G12: Reacted with Nef from different HIV-1 strains (BRU, IIIB, RF, MN). Ovod et al. [1992]

**No.** 1269

MAb ID 13/058

HXB2 Location Nef (60–73)

Author Location Nef (60-73 BH10)

Epitope AQEEEEVGFPVTPQ

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse

References Schneider et al. 1991

• 13/058: Epitope mapped by overlapping decapeptides – core: EEVGFP. Schneider *et al.* [1991]

**No.** 1270

MAb ID 26/028

HXB2 Location Nef (60-73)

**Author Location** Nef (60–73 BH10)

Epitope AQEEEEVGFPVTPQ

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse

References Schneider et al. 1991

 26/028: Epitope mapped by overlapping decapeptides – core: EEVGFPV. Schneider et al. [1991]

**No.** 1271

MAb ID polyclonal

HXB2 Location Nef (61–71)

**Author Location** Nef

Epitope QEEEEVGFPVT

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: Env,

Gag, Nef, Pol

Species (Isotype) rabbit

References Li et al. 2005

Keywords mimics

• In early HIV-1 infection, patients develop autoimmune thrombocytopenia, with Ab directed against beta3 integrin, GPIIIa49-66. Panning with a 7-mer phage display library using rabbit anti-GPIIIa49-66 (CAPESIEFPVSEARVLED), the immunodominant epitope of the identified potential molecular mimicry epitopes with HIV-1 Env (sklFDeGLFn, elfnkTIIFP), Pol (geAPEFPskq), Gag (gktHyMINPl) and Nef (qeeeeVgFPVt, qeeeeVgFPVt, edeGigFPVr, fkLVPVSEae, ssnTPTTNaa) proteins. Pools of these peptides elicted Ab in rabbits that induce platelet oxidation in vitro and thrombocytopenia in vivo upon passive transfer. Nef (qeeeeVgFPVt), Gag (gktHyMINPl), and Nef (fkLVPVSEae) all overlap with known HIV-1 epitopes. Li et al. [2005] (mimics)

**No.** 1272

MAb ID 2E3

HXB2 Location Nef (61–80)

Author Location Nef (61-80 BRU)

Epitope QEEEEVGFPVTPQVPLRPMT

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG1)

References Nilsen et al. 1996; Ovod et al. 1992

- 2E3: There are two MAbs with the name 2E3 the other one binds to integrase. Nilsen *et al.* [1996]
- 2E3: Two isomorphic forms of Nef were identified, 2E3 reacted with the p24 but not p27 form, and was strain specific (MN and BRU reactive, not IIIB or RF). Ovod *et al.* [1992]

No. 1273

MAb ID polyclonal

HXB2 Location Nef (66–97)

Author Location Nef (66–97 LAI)

Epitope VGFPVTPQVPLRPMTYKAAVDLSHFLKEKGGL

Subtype B Neutralizing no Immunogen vaccine

> Vector/Type: lipopeptide Strain: B clade LAI HIV component: Nef Adjuvant:

QS21

Species (Isotype) human (IgG)

References Pialoux et al. 2001

• 28 subjects were vaccinated with six HIV-1 peptides that were selected to be particularly rich in CTL epitopes, presented in lipopeptides with or without adjuvant QS21 – HIV-specific Ab responses were detected in 10/28, proliferative in 11/14, and CTL in 13/24 (54%) of testable volunteers – 10/28 had Ab responses to this peptide (N1), 11/24 had proliferative responses, and CTL responses were detected. Pialoux *et al.* [2001]

**No.** 1274

**MAb ID** F14.11

HXB2 Location Nef (83–88)

**Author Location** Nef (83–88)

**Epitope** AAVDLS

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: Nef

**Species (Isotype)** mouse (IgG2a $\kappa$ )

References Chang et al. 1998; De Santis et al. 1991

- F14.11: Used as a control in a study of Nef-specific single chain Abs constructed from AG11 and EH1. Chang et al. [1998]
- F14.11: The MAb was made to a six an region of Nef that is similar to a region found in thymosin alpha 1 protein the MAb binds to the natural Nef protein. De Santis *et al.* [1991]

No. 1275

**MAb ID** 31/03

HXB2 Location Nef (83-103)

Author Location Nef (82–103 BH10)

**Epitope** AAVDLSHFLKEKGGLEGLIHS

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse

References Schneider et al. 1991

 31/03: Epitope mapped by overlapping decapeptides – mapping suggests complex epitope in this region. Schneider et al. [1991]

**No.** 1276

MAb ID polyclonal

HXB2 Location Nef (90–98)

**Author Location** Nef (NL43)

Epitope FLKEKGGLE

Neutralizing

Immunogen HIV-1 infection, vaccine

Species (Isotype) human, rabbit

References Yamada & Iwamoto 1999

Country Japan

Keywords ADCC, antibody binding site definition and

exposure, antibody generation, complement,

rate of progression

Antibody responses to overlapping 9-mers from the Nef protein were mapped in a set of HIV+ Japanese hemophiliacs. Long term non-progressors among the group were significantly more likely to react to Nef peptide 31 (FLKEKGGLE) (p=0.008). Rabbit polyclonal Abs were raised against this peptide. These Abs bound Nef, could kill infected cells in a complement dependent manner, and the domain near peptide 31 was exposed on the surface of infected T-cells. Yamada & Iwamoto [1999] (ADCC, antibody binding site definition and exposure, antibody generation, complement, rate of progression)

**No.** 1277

MAb ID polyclonal

HXB2 Location Nef (90–98)

**Author Location Nef** 

Epitope FLKEKGGLE

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Yamada et al. 2004

Country Japan

Keywords ADCC, rate of progression

Plasma and PBMC from long term non-progressors can mediate ADCC against Nef infected target cells. Addition of the peptide FLKEKGGLE reduces this activity by half. Patients who were LTNPs were found to make antibodies against this peptide in an earlier study. Anti-Gag antibodies do not elicit ADCC, and Pol proteins are not expressed on the cell surface, in contrast to this Nef epitope. Yamada *et al.* [2004] (ADCC, rate of progression)

**No.** 1278

MAb ID F4

**HXB2 Location** Nef (115–126)

Author Location Nef (115–126 NL-432)

Epitope YHTQGYFPDWQN?

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43

HIV component: Nef

Species (Isotype) mouse (IgG1)

References Otake et al. 1997

Keywords antibody binding site definition and exposure,

antibody generation

• F4: BALB/c mice were immunized with Nef protein to create a series of anti-Nef antibodies. F4 bound to the peptide spanning amino acids 115-126; we inferred the amino acids from the positions in the NL-43 strain. A6 also bound to the complete Nef protein. Otake *et al.* [1997] (antibody binding site definition and exposure, antibody generation)

**No.** 1279 **MAb ID** F2

**HXB2 Location** Nef (115–136)

Author Location Nef (115–137 NL-432)

Epitope YHTQGYFPDWQNYTPGPGVRYP?

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: Nef Adjuvant: Complete

Freund's Adjuvant (CFA)

**Species (Isotype)** mouse (IgG1) **References** Otake *et al.* 1997

Keywords antibody binding site definition and exposure,

antibody generation

• F2: BALB/c mice were immunized with Nef protein to create a series of anti-Nef antibodies. F2 bound to the peptide spanning amino acids 115-137; we inferred the amino acids from the positions in the NL-43 strain. F2 also bound to the complete Nef protein. Otake *et al.* [1997] (antibody binding site definition and exposure, antibody generation)

**No.** 1280

MAb ID polyclonal

HXB2 Location Nef (117-147)

**Author Location** Nef (117–147 LAI)

Epitope TQGYFPDWQNYTPGPGVRYPLTFGWCYKLVP

**Subtype** B **Neutralizing** no **Immunogen** vaccine

Vector/Type: lipopeptide Strain: B clade LAI HIV component: Nef Adjuvant:

OS21

Species (Isotype) human (IgG)

References Pialoux et al. 2001

• 28 subjects were vaccinated with six HIV-1 peptides that were selected to be particularly rich in CTL epitopes, presented in lipopeptides with or without adjuvant QS21 – HIV-specific Ab responses were detected in 25/28, proliferative in 3/24, and CTL in 13/24 (54%) of testable volunteers – 20/28 had antibody responses to this particular peptide (N2), 3/24 had proliferative responses, and CTL responses were detected. Pialoux *et al.* [2001]

**No.** 1281

MAb ID polyclonal

HXB2 Location Nef (118-133)

**Author Location** Nef (118–133)

Epitope QGYFPDWQNYTPGPGV

Neutralizing no

Immunogen vaccine

Vector/Type: DNA Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG)

References Tahtinen et al. 2001

• BALB/c mice were immunized with a pBN-vector expressing HIV-1 nef, rev, or tat genes—DNA loaded onto gold microparticles was delivered using a gene gun, and DNA dissolved in saline was given intradermally or intramuscularly—Nef gene gun immunized mice showed the strongest and most long-lasting (6 months) Ab, CTL and proliferative responses—the highest IgG1/IgG2a ratio was observed in the gene gun immunized mice—three Ab binding sites were found in Nef using peptide mapping, although some sera reacted only to complete Nef—Rev- or Tat-immunized mice did not generate an Ab response. Tahtinen *et al.* [2001]

**No.** 1282

MAb ID polyclonal

**HXB2 Location** Nef (119–168)

Author Location Nef (118–167 LAI, BRU)

Epitope GYFPDWQNYTPGPGVRYPLTFGWCYKLVPVEP-

**DKVEEANKGENTSLLHPV** 

Subtype B

Neutralizing

Immunogen HIV-1 infection, vaccine

Vector/Type: protein, PLG microparticle Strain: B clade BRU, B clade LAI HIV component: Nef Adjuvant: Complete Freund's

Adjuvant (CFA), PLG

Species (Isotype) mouse (IgG1)

References Maksiutov et al. 2002; Moureau et al. 2002

- This epitope is similar to a fragment of the human protein Bonederived growth factor, PLEPAKLEE, and to Hematopoietic progenitor cell antigen CD34, TSLHPVSQHG. Maksiutov *et al.* [2002]
- Nef encapsulated in poly(DL-lactide-co-glycolide) (PLG) had a more prolonged Ab response than Nef in PBS or in Freund's adjuvant (CFA), still strong after 7 months the response was predominantly IgG1, a Th2 immune response three linear epitopes, Nef 32-64, 118-167, and 185-205, were frequently recognized by the sera of mice immunized with NefPLG or Nef-CFA, but not after immunization with Nef in PBS, which seemed to preferentially stimulate an Ab response to conformational epitopes. Moureau et al. [2002]

**No.** 1283

MAb ID F3

**HXB2 Location** Nef (128–137)

Author Location Nef (128–137 NL-432)

Epitope TPGPGVRYPL?

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: Nef Adjuvant: Complete

Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgG1)

References Kawai et al. 2003; Otake et al. 1997

Keywords antibody binding site definition and exposure,

antibody generation, complement

- F3: Used as a control for Nef binding in a study designed to elicit anti-Nef IgM human Abs in trans-chromosome mice that carry human chromosomes responsible for Ig production. These mice were immunized with recombinant Nef protein. Human heavy chain, mouse light chain anti-Nef IgM were obtained. The hope was that these human IgM anti-Nef chimeric Abs would activate complement mediated cytolysis; C3 deposition was induced but not at high enough levels to induce lysis. Kawai et al. [2003] (complement)
- F3: BALB/c mice were immunized with Nef protein to create a series of anti-Nef antibodies. F3 bound to the peptide spanning amino acids 128-137; we inferred the amino acids from the positions in the NL-43 strain. F3 also bound to the complete Nef protein. Otake *et al.* [1997] (antibody binding site definition and exposure, antibody generation)

**No.** 1284

MAb ID F8

HXB2 Location Nef (128-137)

Author Location Nef (128-137 NL-432)

Epitope TPGPGVRYPL?

Subtype B

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA)

**Species (Isotype)** mouse (IgM) **References** Otake *et al.* 1997

**Keywords** antibody binding site definition and exposure, antibody generation

• F8: BALB/c mice were immunized with Nef protein to create a series of anti-Nef antibodies. F8 bound to the peptide spanning amino acids 128-137; we inferred the amino acids from the positions in the NL-43 strain. F8 also bound to the complete Nef protein. Otake *et al.* [1997] (antibody binding site definition and exposure, antibody generation)

No. 1285

MAb ID polyclonal

**HXB2 Location** Nef (143–151)

**Author Location** Nef

**Epitope** FKLVPVSEAE

Neutralizing

Immunogen vaccine

Vector/Type: peptide HIV component: Env, Gag, Nef, Pol

Species (Isotype) rabbit

References Li et al. 2005

Keywords mimics

• In early HIV-1 infection, patients develop autoimmune throm-bocytopenia, with Ab directed against beta3 integrin, GPIIIa49-66. Panning with a 7-mer phage display library using rabbit anti-GPIIIa49-66 (CAPESIEFPVSEARVLED), the immunodominant epitope of the identified potential molecular mimicry epitopes with HIV-1 Env (skIFDeGLFn, elfnkTIIFP), Pol (geAPEFPskq), Gag (gktHyMINPl) and Nef (qeeeeVgFPVt, qeeeeVgFPVt, edeGigFPVr, fkLVPVSEae, ssnTPTTNaa) proteins. Pools of these peptides elicted Ab in rabbits that induce platelet oxidation in vitro and thrombocytopenia in vivo upon

passive transfer. Nef (qeeeeVgFPVt), Gag (gktHyMINPl), and Nef (fkLVPVSEae) all overlap with known HIV-1 epitopes. Li *et al.* [2005] (mimics)

**No.** 1286

MAb ID F1

**HXB2 Location** Nef (148–157)

Author Location Nef (148–157 IIIB)

Epitope VEPDKVEEAN

Neutralizing

**Immunogen** 

Species (Isotype) mouse (IgM)

**References** Haynes *et al.* 2005; Fujii *et al.* 1996b; Fujii *et al.* 1996c; Otake *et al.* 1994; Fujii *et al.* 

1993

- F1: There is a Nef (Fujii1993) and a CD4BS (Haynes2005)
   MAb that are called F1. Fujii et al. [1993]; Haynes et al. [2005]
- F1: Insect cells expressing myristylated Nef proteins on their cell surface can induce cytolysis of unstimulated CD4+ cells this response is not due to MHC restricted CTL activity the cell surface of Nef expressing insect cells carry Nef that can be recognized by MAbs E7 and E9 but not F1. Fujii *et al.* [1996c]
- F1: A carboxy-terminal domain of Nef on the cell surface induces cytolysis of CD4+ T cells. Fujii *et al.* [1996b]
- F1: The C-term end of Nef is accessible to Abs at the cell surface stained IIIB/M10, but not MN/M10, cells. Fujii *et al.* [1993]; Otake *et al.* [1994]

**No.** 1287

MAb ID 2F2

**HXB2 Location** Nef (151–170)

**Author Location** Nef (151–170 BRU)

Epitope DKVEEANKGENTSLLHPVSL

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: Nef

Species (Isotype) mouse (IgG1)

**References** Maksiutov *et al.* 2002; Ranki *et al.* 1995; Saito *et al.* 1994; Ovod *et al.* 1992

- 2F2: UK Medical Research Council AIDS reagent: EVA3067.3.
- 2F2: This epitope is similar to a fragment of the human protein Hematopoietic progenitor cell antigen CD34, TSLHPVSQHG. Maksiutov et al. [2002]
- 2F2: One of four antibodies used in combination to show HIV Nef protein expressed in astrocytes from 7/14 brain samples from HIV+ individuals – Nef expression associated with dementia. Ranki et al. [1995]
- 2F2: Over-expression of Nef in astrocytes from postmortem pediatric CNS tissue. Saito et al. [1994]
- 2F2: Strain specific (MN and BRU reactive, not IIIB or RF).
   Ovod et al. [1992]

No. 1288

MAb ID E9

HXB2 Location Nef (158–181)

**Author Location** Nef (158–206 IIIB)

Epitope KGENTSLLHPVSLHGMDDPEREVL

Neutralizing

Immunogen

Species (Isotype) mouse (IgM)

**References** Maksiutov *et al.* 2002; Fujii *et al.* 1996b; Fujii *et al.* 1996c; Otake *et al.* 1994; Fujii *et al.* 1993

 E9: This epitope is similar to a fragment of the human protein Hematopoietic progenitor cell antigen CD34, TSLHPVSQHG. Maksiutov et al. [2002]

- E9: A carboxy-terminal domain of Nef on the cell surface induces cytolysis of CD4+ T cells. Fujii *et al.* [1996b]
- E9: Insect cells expressing myristylated Nef proteins on their cell surface can induce cytolysis of unstimulated CD4+ cells this response is not due to MHC restricted CTL activity the cell surface of Nef expressing insect cells carry Nef that can be recognized by MAbs E7 and E9 but not F1. Fujii *et al.* [1996c]
- E9: The C-term end of Nef is accessible to Abs at the cell surface – stained IIIB/M10, but not MN/M10, cells. Fujii et al. [1993]; Otake et al. [1994]

**No.** 1289

MAb ID 3E6

**HXB2 Location** Nef (161–180) **Author Location** Nef (161–180 BRU)

Epitope NTSLLHPVSLHGMDDPEREV

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU HIV component: Nef

Species (Isotype) mouse (IgG1)

**References** Maksiutov *et al.* 2002; Ranki *et al.* 1995; Saito *et al.* 1994; Ovod *et al.* 1992

- 3E6: UK Medical Research Council AIDS reagent: EVA3067.4.
- 3E6: This epitope is similar to a fragment of the human protein Hematopoietic progenitor cell antigen CD34, TSLHPVSQHG. Maksiutov *et al.* [2002]
- 3E6: Faintly cross-reactive with astrocytes of uninfected control samples. Ranki *et al.* [1995]
- 3E6: Reacted with Nef from different HIV-1 strains (BRU, IIIB, RF, MN). Ovod *et al.* [1992]

**No.** 1290

MAb ID E5

**HXB2 Location** Nef (170–181)

**Author Location** Nef (170–181)

Epitope LHGMDDPEREVL?

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade NL43 HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA)

Species (Isotype) mouse (IgM)

References Otake et al. 1997

**Keywords** antibody binding site definition and exposure, antibody generation

• E5: BALB/c mice were immunized with Nef protein to create a series of anti-Nef antibodies. E5 bound to the peptide spanning amino acids 170-181; we inferred the amino acids from the positions in the NL-43 strain. E5 also bound to the complete Nef protein. Otake *et al.* [1997] (antibody binding site definition and exposure, antibody generation)

**No.** 1291

MAb ID 2A3

**HXB2 Location** Nef (171–190)

Author Location Nef (171–190 BRU)

Epitope HGMDDPEREVLEWRFDSRLA

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG1)

References Ovod et al. 1992

 2A3: Reacted with Nef from different HIV-1 strains (BRU, IIIB, MN, but not RF). Ovod et al. [1992]

No. 1292

MAb ID 2E4

**HXB2 Location** Nef (171–190)

**Author Location** Nef (171–190 BRU)

Epitope HGMDDPEREVLEWRFDSRLA

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG1)

References Ovod et al. 1992

 2EA: Reacted with Nef from different HIV-1 strains (BRU, IIIB, MN but not RF). Ovod et al. [1992]

No. 1293

MAb ID 2H12

**HXB2 Location** Nef (171–190)

Author Location Nef (171–190 BRU)

Epitope HGMDDPEREVLEWRFDSRLA

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG1)

**References** Ranki *et al.* 1995; Saito *et al.* 1994; Ovod *et al.* 1992

- 2H12: One of four antibodies used in combination to show HIV Nef protein expressed in astrocytes from 7/14 brain samples from HIV+ individuals – Nef expression associated with dementia. Ranki et al. [1995]
- 2H12: Over-expression of Nef in astrocytes from postmortem pediatric CNS tissue. Saito et al. [1994]
- 2H12: Reacted with Nef from different HIV-1 strains (BRU, IIIB, RF, MN). Ovod *et al.* [1992]

**No.** 1294

MAb ID 3A2

**HXB2 Location** Nef (171–190)

**Author Location** Nef (171–190 BRU)

Epitope HGMDDPEREVLEWRFDSRLA

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade BRU

HIV component: Nef

Species (Isotype) mouse (IgG1)

Nef Antibodies HIV Antibodies Tables

**References** Ranki *et al.* 1995; Saito *et al.* 1994; Ovod *et al.* 1992

- 3A2: UK Medical Research Council AIDS reagent: EVA3067.5.
- 3A2: One of four antibodies used in combination to show HIV Nef protein expressed in astrocytes from 7/14 brain samples from HIV+ individuals – Nef expression associated with dementia. Ranki *et al.* [1995]
- 3A2: Over-expression of Nef in astrocytes from postmortem pediatric CNS tissue. Saito *et al.* [1994]
- 3A2: Reacted with Nef from different HIV-1 strains (BRU, IIIB, RF, MN). Ovod *et al.* [1992]

**No.** 1295

MAb ID NF1A1

**HXB2 Location** Nef (173–206)

Author Location Nef (173–206)

 ${\bf Epitope} \ \ {\tt MDDPEREVLEWRFDSRLAFHHVARELHPEYFK-}$ 

NC

**Neutralizing** 

Immunogen

Species (Isotype) mouse

References Kaminchik et al. 1990

 NF1A1: Recognizes the Nef protein of the two isolates BH10 and LAV1 – low affinity. Kaminchik et al. [1990]

**No.** 1296

MAb ID polyclonal

**HXB2 Location** Nef (186–206)

Author Location Nef (185–205 LAI, BRU)

Epitope DSRLAFHHVARELHPEYFKNC

Subtype B

Neutralizing

Immunogen HIV-1 infection, vaccine

Vector/Type: protein, PLG microparticle Strain: B clade BRU, B clade LAI HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA), PLG

Species (Isotype) mouse (IgG1)

References Moureau et al. 2002

• Nef encapsulated in poly(DL-lactide-co-glycolide) (PLG) had a more prolonged Ab response than Nef in PBS or in Freund's adjuvant (CFA), still strong after 7 months – the response was predominantly IgG1, a Th2 immune response – three linear epitopes, Nef 32-64, 118-167, and 185-205, were frequently recognized by the sera of mice immunized with NefPLG or Nef-CFA, but not after immunization with Nef in PBS, which seemed to preferentially stimulate an Ab response to conformational epitopes. Moureau et al. [2002]

**No.** 1297

MAb ID E7

**HXB2 Location** Nef (192–206)

Author Location Nef (192-206 IIIB)

Epitope HHVARELHPEYFKNC

Neutralizing

Immunogen

Species (Isotype) mouse (IgM)

**References** Fujii *et al.* 1996d; Fujii *et al.* 1996b; Fujii *et al.* 1996a; Fujii *et al.* 1996c; Otake *et al.* 1994; Fujii *et al.* 1993

- E7: Insect cells expressing myristylated Nef proteins on their cell surface can induce cytolysis of unstimulated CD4+ cells this response is not due to MHC restricted CTL activity the cell surface of Nef expressing insect cells carry Nef that can be recognized by MAbs E7 and E9 but not F1. Fujii *et al.* [1996c]
- E7: Nef forms a homomeric oligomerizing structure, and using E7 and membrane immunofluorescence or immunoelectron microscopy, was shown to clusters on the surface of HIV-1 infected CD4+ cells. Fujii *et al.* [1996a]
- E7: A carboxy-terminal domain of Nef on the cell surface induces cytolysis of CD4+ T cells. Fujii *et al.* [1996b]
- E7: Soluble Nef inhibits proliferation of CD4+ cells, and Nef cross-linking by MAbs may induce anti-CD4 cytocidal activity sera from HIV+ individuals contain soluble Nef, thus this may be important for immune dysfunction and disease progression. Fujii *et al.* [1996d]
- E7: The C-term end of Nef is accessible to Abs at the cell surface stained IIIB/M10, but not MN/M10, cells. Fujii *et al.* [1993]; Otake *et al.* [1994]

**No.** 1298

MAb ID AE6

**HXB2 Location** Nef (194–206)

Author Location Nef (LAI)

**Epitope** VARELHPEYFKNC

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type C-term

Vector/Type: protein, PLG microparticle Research Contact Frank Jirik, Centre for Molecular Med and

Therapeutics, U. B. C., Vancouver, B. C.

Canada

References Chang et al. 1998

AE6: The light and heavy chains of three MAbs (AG11, AE6, EH1) specific to C-terminus of NEF were cloned and variable regions sequenced – the complementarity determining regions (CDR) of AG11 and AE6 were highly related (95.1% at the DNA level) and bound LAI Nef, but not SF2 Nef – EH1 bound to SF2 and LAI and cross-competed AG11 and AE6 but had a distinctive CDR (57.9% similar to AG11) – single chain Abs were constructed from AG11 and EH1. Chang et al. [1998]

**No.** 1299

MAb ID AG11

**HXB2 Location** Nef (194–206)

**Author Location** Nef (LAI)

**Epitope** VARELHPEYFKNC

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type C-term

HIV Antibodies Tables Nef Antibodies

Research Contact Frank Jirik, Centre for Molecular Med and Therapeutics, U. B. C., Vancouver, B. C.

Canada

- 3B4B: The goal of this study was to elicit anti-Nef IgM human Abs in trans-chromosome mice that carry human chromosomes responsible for Ig production. These mice were immunized

References Chang et al. 1998

• AG11: The light and heavy chains of three MAbs (AG11, AE6, EH1) specific to C-terminus of NEF were cloned and variable regions sequenced – the complementarity determining regions (CDR) of AG11 and AE6 were highly related (95.1% at the DNA level) and bound LAI Nef, but not SF2 Nef – EH1 bound to SF2 and LAI and cross-competed AG11 and AE6 but had a distinctive CDR (57.9% similar to AG11) – single chain Abs were constructed from AG11 and EH1 and subcloned into a eukaryotic expression vector with a green fluorescent protein marker to allow intracellular expression – the single chain Abs bind Nef intracellularly and may be useful to better understand the role of Nef and as a gene therapy model. Chang *et al.* [1998]

**No.** 1300

MAb ID EH1

**HXB2 Location** Nef (194–206) **Author Location** Nef (SF2)

Epitope MARELHPEYYKDC

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef

**Species (Isotype)** mouse (IgG1 $\kappa$ )

Ab Type C-term

Research Contact Frank Jirik, Centre for Molecular Med and Therapeutics, U. B. C., Vancouver, B. C. Canada

References Chang et al. 1998

• EH1: The light and heavy chains of three MAbs (AG11, AE6, EH1) specific to C-terminus of NEF were cloned and variable regions sequenced – the complementarity determining regions (CDR) of AG11 and AE6 were highly related (95.1% at the DNA level) and bound LAI Nef, but not SF2 Nef – EH1 bound to SF2 and LAI and cross-competed AG11 and AE6 but had a distinctive CDR (57.9% similar to AG11) – single chain Abs were constructed from AG11 and EH1 and subcloned into a eukaryotic expression vector with a green fluorescent protein marker to allow intracellular expression – the single chain Abs bind Nef intracellularly and may be useful to better understand the role of Nef and as a gene therapy model. Chang *et al.* [1998]

**No.** 1301

MAb ID 3B4B

**HXB2 Location** Nef

Author Location Nef Epitope

Neutralizing

Immunogen vaccine

Vector/Type: protein HIV component: Nef Adjuvant: Incomplete Freund's Adjuvant

(IFA)

Species (Isotype) transgenic mouse (IgM)

References Kawai et al. 2003

Keywords antibody generation, complement

• 3B4B: The goal of this study was to elicit anti-Nef IgM human Abs in trans-chromosome mice that carry human chromosomes responsible for Ig production. These mice were immunized with recombinant Nef protein. Two human heavy chain, mouse light chain anti-Nef IgM were obtained, 3B4B and 3H3E; 3B4B was able to stain MOLT4/IIIB cells with greater intensity. The hope was that these human IgM anti-Nef chimeric Abs would activate complement mediated cytolysis; C3 deposition was induced but not at high enough levels to induce lysis. Kawai *et al.* [2003] (antibody generation, complement)

**No.** 1302

MAb ID 3H3E

**HXB2 Location** Nef

**Author Location** Nef

**Epitope** 

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein HIV component: Nef Adjuvant: Complete Freund's Adjuvant (CFA)

Species (Isotype) transgenic mouse (IgM)

References Kawai et al. 2003

Keywords antibody generation, complement

• 3H3E: The goal of this study was to elicit anti-Nef IgM human Abs in trans-chromosome mice that carry human chromosomes responsible for Ig production. These mice were immunized with recombinant Nef protein. Two human heavy chain, mouse light chain anti-Nef IgM were obtained, 3B4B and 3H3E; 3B4B was able to stain MOLT4/IIIB cells with greater intensity. The hope was that these human IgM anti-Nef chimeric Abs would activate complement mediated cytolysis; C3 deposition was induced but not at high enough levels to induce lysis. Kawai et al. [2003] (antibody generation, complement)

**No.** 1303

**MAb ID** 6.1

**HXB2 Location** Nef

**Author Location** Nef (JRCSF)

**Epitope** 

Subtype B

Neutralizing

Immunogen

Species (Isotype) mouse

References Ranki et al. 1995

- 6.1: Raised against CNS primary isolates, stains astrocytes more densely than other Nef MAbs – Nef expression associated with dementia. Ranki *et al.* [1995]
- 6.1: NIAID Repository number 1123. Ranki et al. [1995]

**No.** 1304

MAb ID NF2B2

HXB2 Location Nef

Author Location Nef (20–78 BH10)

Epitope

**Neutralizing** 

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: Nef

Species (Isotype) mouse

HIV-1 Antibodies HIV Antibodies Tables

References Kaminchik et al. 1990

 NF2B2: NIH AIDS Research and Reference Reagent Program: 456.

• NF2B2: Recognizes the Nef protein of the two isolates BH10 and LAV1. Kaminchik *et al.* [1990]

**No.** 1305

MAb ID NF3A3

**HXB2 Location** Nef

Author Location Nef (20-78 BH10)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: Nef

Species (Isotype) mouse

References Kaminchik et al. 1990

• NF3A3: Recognizes the Nef protein of the two isolates BH10 and LAV1 – low affinity. Kaminchik *et al.* [1990]

**No.** 1306

MAb ID NF8B4

**HXB2 Location** Nef

Author Location Nef (BH10)

**Epitope** 

Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade BH10

HIV component: Nef

Species (Isotype) mouse

References Kaminchik et al. 1990

 NF8B4: Does not recognize Nef CNBr cleavage products – recognizes intact BH10 Nef but not LAV1 Nef. Kaminchik et al. [1990]

**No.** 1307

MAb ID polyclonal

**HXB2 Location** Nef

**Author Location** Nef

Epitope

Subtype B Neutralizing

Immunogen vaccine

Vector/Type: protein Strain: B clade LAI,

SIV *HIV component:* gp120, Nef, Tat *Adjuvant:* AS02A (oil-in-water emulsion, 3D-monophosphoryl lipid A, QS21), AS06 (CpG,

aluminum hydroxide)

Species (Isotype) macaque (IgG)

References Voss et al. 2003

**Keywords** adjuvant comparison, variant crossrecognition or cross-neutralization

 Monkeys were immunized with different combinations of gp120/NefTat/SIV-Tat three times over a three-month time period and intravenously challenged with SHIV 89.6P four weeks after the last immunzation. Vaccination induced anti-Tat, -Nef and -gp120 Ab responses that decreased until challenge. Neutralizing Ab responses against HIV-1 MN and HIV-1 W61D were observed. Vaccination with the combination of all three proteins reduced viral load and kept animals from a CD4+ T cell decline, preventing AIDS for more than 2.5 years. The adjuvant AS02A elicited NAbs and protected from disease, while AS06 did not. gp120 alone or TatNef/SIV-Tat without gp120 could not protect from disease. Voss *et al.* [2003] (adjuvant comparison, variant cross-recognition or cross-neutralization)

**No.** 1308

MAb ID AE6

**HXB2 Location** Nef

**Author Location Nef** 

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) mouse

Ab Type C-term

Research Contact James Hoxie, Div of AIDS, NIAID, NIH

References Tornatore et al. 1994; Greenway et al. 1994

AE6: NIH AIDS Research and Reference Reagent Program:

#### IV-C-19 HIV-1 Antibodies

**No.** 1309

MAb ID

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype)

References Goepfert 2003

Keywords review

 A general review of anti-HIV human immune responses and the implications of these responses for vaccines, summarizing neutralizing antibodies, CD4+ and CD8+ T cell responses. A general overview of methods used to study these responses is presented. Goepfert [2003] (review)

**No.** 1310

MAb ID

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen vaccine

*Adjuvant:* CD40, CD80, CD86, Complete Freund's Adjuvant (CFA), GM-CSF, IFN $\gamma$ , IL-12, IL-15, IL-18, IL-1 $\alpha$ , IL-2, IL-2/Ig, IL-4, IL-7, CpG immunostimulatory sequence (ISS), Tumor Necrosis Factor  $\alpha$  (TNF $\alpha$ ), Tumor Necrosis Factor  $\beta$  (TNF $\beta$ ), M-CSF, IL-8, RANTES

Species (Isotype)

References Mitchison & Sattentau 2005

Keywords adjuvant comparison, review, Th1, Th2

HIV-1 Antibodies Tables HIV-1 Antibodies

• Review summarizes mechanisms of immunoregulation relevant for new vaccine development, with a brief summary of adjuvant triggering innate immunity through Toll-like receptors (TLRs), Nod molecules, and other activators. DNA encoded adjuvants that have been tested in DNA vaccines are summarized. The balance between Th1 (CTL activating) and Th2 (B cell activating) responses is discussed, and it is noted that BALB/c mice are predominately Th2 responders, C57BL Th1. Mitchison & Sattentau [2005] (adjuvant comparison, Th1, Th2, review)

**No.** 1311

MAb ID

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

**Neutralizing** 

**Immunogen** 

Species (Isotype) human

References Piontkivska & Hughes 2006

Keywords escape

• The greatest amino acid diversity is found in sites in the HIV genome that are spanned by antibody epitopes. Sites spanned by CTL epitopes, not but not by antibody epitopes, showed reduced amino acid diversity even in comparison to non-epitope sites. However, mutations within CTL epitopes were more likely to be convergent than mutations within antibody epitopes. These patterns were consistent both in Gag and Env. Piontkivska & Hughes [2006] (escape)

**No.** 1312

**MAb ID** 1.4C

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 adjacent to CD4BS

References Haynes et al. 2005

Keywords antibody binding site definition and exposure

1.4C: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 1.4C has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

**No.** 1313

MAb ID 1.4G

**HXB2 Location** HIV-1

**Author Location** 

Epitope

Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 adjacent to CD4BS

References Haynes et al. 2005

Keywords antibody binding site definition and exposure

• 1.4G: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. Haynes *et al.* [2005] (antibody binding site definition and exposure)

**No.** 1314

**MAb ID** 1.9E

**HXB2 Location** HIV-1

**Author Location** 

Epitope

Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 CCR5BS

References Haynes et al. 2005

Keywords antibody binding site definition and exposure

• 1.9E: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 1.9E has no indication of polyspecific autoreactivity. Haynes *et al.* [2005] (antibody binding site definition and exposure)

**No.** 1315

**MAb ID** 1.9F

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype) human

Ab Type gp120 CCR5BS

References Haynes et al. 2005

Keywords antibody binding site definition and exposure

• 1.9F: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 1.9F has no indication of polyspecific autoreactivity. Haynes *et al.* [2005] (antibody binding site definition and exposure)

**No.** 1316

**MAb ID** 12.19

HXB2 Location HIV-1

Author Location

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) (IgG)

Ab Type gp120 V3

References Koefoed et al. 2005

HIV-1 Antibodies HIV Antibodies Tables

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, kinetics

• 12.19: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. This antibody bound to a V3-fusion protein. Koefoed *et al.* [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

**No.** 1317

**MAb ID** 12.9

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) (IgG)

Ab Type gp120 V3

References Koefoed et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, kinetics

• 12.9: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. This antibody bound to a V3-fusion protein. Koefoed *et al.* [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

**No.** 1318

**MAb ID** 13a15

**HXB2** Location HIV-1

Author Location (JRFL)

Epitope

Subtype B

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

Ab Type gp120 CD4BS

References Koefoed et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, kinetics

• 13a15: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. Most of the

Fabs bound to the CD4BS, including this one, as they could be blocked with sCD4 and murine anti-CD4BS Abs. Fab 13a15 was not neutralizing. Koefoed *et al.* [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

**No.** 1319

**MAb ID** 13a23

HXB2 Location HIV-1

**Author Location (JRFL)** 

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

Ab Type gp120 CD4BS

References Koefoed et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, kinetics

• 13a23: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. Most of the Fabs bound to the CD4BS, including this one, as they could be blocked with sCD4 and murine anti-CD4BS Abs; 13a23 was somewhat different from the other CD4BS Fabs isolated in this study, in that its binding was enhanced by anti-C1 MAbs. Fab 13a23 was not neutralizing. Koefoed *et al.* [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

**No.** 1320

**MAb ID** 13a3

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

Ab Type gp120 CD4BS

References Koefoed et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, vari-

able domain, kinetics

• 13a3: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. Most of the Fabs bound to the CD4BS, including this one, as they could be blocked with sCD4 and murine anti-CD4BS Abs. Fab 13a3 weakly neutralized MN, but not HXB2 Ba-L or JRFL. Koefoed et al. [2005] (antibody binding site definition and exposure,

**HIV Antibodies Tables HIV-1 Antibodies** 

antibody generation, kinetics, antibody sequence, variable Author Location (LAI) domain)

No. 1321 **MAb ID** 13a6 **HXB2** Location HIV-1

**Author Location** 

**Epitope** Subtype B Neutralizing no

Immunogen HIV-1 infection Species (Isotype) human (IgG) Ab Type gp120 CD4BS

References Koefoed et al. 2005

Keywords antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, kinetics

• 13a6: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to using bone marrow for generating libraries, which is often difficult to obtain. Most of the Fabs bound to the CD4BS, including this one, as they could be blocked with sCD4 and murine anti-CD4BS Abs. Fab 13a6 was not neutralizing. Koefoed et al. [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

No. 1322 **MAb ID** 13a7 **HXB2** Location HIV-1 **Author Location Epitope** Subtype B Neutralizing no Immunogen HIV-1 infection Species (Isotype) human (IgG) Ab Type gp120 CD4BS

> References Koefoed et al. 2005 **Keywords** antibody binding site definition and exposure,

> antibody generation, antibody sequence, variable domain, kinetics

• 13a7: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. Most of the Fabs bound to the CD4BS, including this one, as they could be blocked with sCD4 and murine anti-CD4BS Abs. Fab 13a7 was not neutralizing. Koefoed et al. [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

No. 1323 **MAb ID** 13b18 **HXB2** Location HIV-1

**Epitope** Subtype B Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG) Ab Type gp120 CD4BS

References Koefoed et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, kinetics

• 13b18: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. Most of the Fabs bound to the CD4BS, including this one, as they could be blocked with sCD4 and murine anti-CD4BS Abs. Fab 13b18 was not neutralizing. Koefoed et al. [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

**No.** 1324

**MAb ID** 13b23

**HXB2 Location** HIV-1

**Author Location (LAI)** 

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

Ab Type gp120 C1

References Koefoed et al. 2005

Keywords antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, kinetics

• 13b120: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. Most of the Fabs bound to the CD4BS, but not 13b120; this Fab was partially inhibited by anti-C1 mAb MAG45, and enhanced by CD4i MAb b17 and anti-C1 MAb 1331290. Koefoed et al. [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

**No.** 1325

**MAb ID** 13b53

HXB2 Location HIV-1

**Author Location (LAI)** 

**Epitope** 

Subtype B

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

HIV-1 Antibodies HIV Antibodies Tables

**Ab Type** gp120 CD4BS **References** Koefoed *et al.* 2005

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, variable domain, kinetics

• 13b53: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after preselection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. Most of the Fabs bound to the CD4BS, including this one, as they could be blocked with sCD4 and murine anti-CD4BS Abs. Fab 13b53 was not neutralizing. Koefoed *et al.* [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

**No.** 1326

**MAb ID** 13b61

**HXB2 Location** HIV-1

**Author Location** (LAI)

**Epitope** 

Subtype B

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

Ab Type gp120 CD4BS

References Koefoed et al. 2005

**Keywords** antibody binding site definition and exposure, antibody generation, antibody sequence, vari-

able domain, kinetics

• 13b61: IgG antibody phage display libraries were created from HIV-1 + individuals using unselected PBMC or after pre-selection of PBMC with gp120 using affinity columns. This was one of 2 gp120 Abs from the unselected library; 9 Abs were were obtained from a 10-fold smaller pre-selected library. This approach offers an alternative to generating libraries from bone marrow, which is often difficult to obtain. Most of the Fabs bound to the CD4BS, including this one, as they could be blocked with sCD4 and murine anti-CD4BS Abs. Fab 13b61 could neutralize HXB2 at 25 ug/ml, but not MN, Ba-L or JRFL. Koefoed *et al.* [2005] (antibody binding site definition and exposure, antibody generation, kinetics, antibody sequence, variable domain)

No. 1327

**MAb ID** 2.5E

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 CCR5BS

References Haynes et al. 2005

**Keywords** antibody binding site definition and exposure

2.5E: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 2.5E has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

**No.** 1328

**MAb ID** 2191

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype)

Ab Type gp120 V3

Research Contact Abraham Pinter, Public Health Research Institute, Newark, NJ. pinter@phri.org

References Pinter et al. 2005

Keywords antibody binding site definition and exposure

• 2191: This study is about the V2 MAb C108g, that is type-specific and neutralizes BaL and HXB2. JR-FL is a neutralization resistant strain; modification of JRFL at V2 positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The modification in V2 also increased neutralization sensitivity to V3 MABs 4117c, 2219, 2191, and 447-52D, but only had minor effects on neutralization by CD4BS MAb 5145A, and broadly neutralizing MAbs IgG1b12, 2G12, and 2F5. Pinter *et al.* [2005] (antibody binding site definition and exposure)

**No.** 1329

MAb ID 25G

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human

**Ab Type** gp120 CD4BS

References Haynes et al. 2005

**Keywords** antibody binding site definition and exposure 25G: Of 35 Env-specific MAbs tested, only 2F5, 4E10,

IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 25G has no indication of polyspecific autoreactivity. Haynes *et al.* [2005] (antibody binding site

definition and exposure)

**No.** 1330

**MAb ID** 2601

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype A, CRF02\_AG

Neutralizing

HIV-1 Antibodies Tables HIV-1 Antibodies

Immunogen HIV-1 infection

**Species (Isotype)** human **Ab Type** gp120 V3

References Krachmarov et al. 2005

**Keywords** antibody binding site definition and exposure, subtype comparisons, variant cross-recognition or cross-neutralization

• 2601: Sera from subtype A infected individuals from Cameroon have antibodies that react strongly with subtype A and subtype B V3 loops in fusion proteins, and neutralize SF162 pseudotypes, while sera from 47 subtype B infected individuals reacted only with subtype B V3s. Sera from Cameroon did not neutralize primary A or B isolates, due to indirect masking by the V1/V2 domain rather than due to loss of the target epitope. 2601 was derived from a person infected with a clade A or CRF02 virus, and binds to A but not to B V3 loops. Neutralization of JR-FL and SF162(UG V3) by anti-V3 MAbs 2557, 2558, 2601, but not subtype A primary isolates despite binding to the subtype A V3 loops, suggested masking by V1V2 blocking of neutralization by these antibodies. Krachmarov et al. [2005] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 1331

**MAb ID** 2909

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Subtype B

Neutralizing

Immunogen HIV-1 infection

**Species** (**Isotype**) human ( $IgG1\lambda$ )

Ab Type quaternary structure

Research Contact Susan Zolla-Pazner (Zol-

las01@mcrcr6.med.nyu), NYU, NY

References Gorny et al. 2005

**Keywords** antibody generation, antibody sequence, variable domain, subtype comparisons, variant cross-recognition or cross-neutralization

• 2909: 2909 is a NAb that was produced by fusion of heteromyeloma SHM-D33 with Epstein-Barr virus transformed PBMC and selection by a neutralization assay. The PBMC were derived from an HIV-1 infected individual who maintained a low viral load after 15 years of infection with no therapy. The MAb very potently neutralizes SF162, but has a narrow range of activity, and did not neutralize autologous virus, nor primary isolates from clade A (VI191, CA1, and 92RW021), clades B (BX08, CA5, and BaL), clade C (95ZW2036) and clade F (CA20 and 93BR029). Sequence analysis of the variable domain of the heavy chain of 2909 shows that it is comprised of IgHV3-43, IgHJ6, and IgHD5-12. 2909 recongizes a quartenary structure present on intact SF162 virions and does not bind to soluble or recombinant Envelope proteins. ELISAbased competition assays and subsequent mutational analysis determined that the CD4BS and V2 and V3 loops contribute to the 2909 epitope: 2909 binding was inhibited by MAbs 447-52d (anti-V3), 830A (anti-V2), and IgG1b12 (anti-CD4BS) and sCD4. 2909 was not inhibited by MAbs 670 (anti-C5), 1418 (irrelevant control MAb), nor 2G12 (anti-carbohydrate); in fact, 2G12 enhanced 2909 binding. Gorny *et al.* [2005] (antibody generation, variant cross-recognition or cross-neutralization, subtype comparisons, antibody sequence, variable domain)

**No.** 1332

**MAb ID** 4.11C

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype) human

Ab Type gp120 adjacent to CD4BS

References Haynes et al. 2005

**Keywords** antibody binding site definition and exposure • 4.11C: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 4.11C has no indication of polyspecific autoreactivity. Haynes *et al.* [2005] (**antibody binding site** 

No. 1333

**MAb ID** 4.6H

definition and exposure)

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 adjacent to CD4BS

References Haynes et al. 2005

**Keywords** antibody binding site definition and exposure • 4.6H: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 4.6H has no indication of polyspecific autoreactivity. Haynes *et al.* [2005] (antibody binding site definition and exposure)

**No.** 1334

**MAb ID** 4.8E

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 CCR5BS

**References** Haynes et al. 2005

Keywords antibody binding site definition and exposure

**HIV-1 Antibodies HIV Antibodies Tables** 

• 4.8E: Of 35 Env-specific MAbs tested, only 2F5, 4E10, Author Location IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 4.8E has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

**No.** 1335 **MAb ID** 5145A **HXB2 Location** HIV-1 **Author Location Epitope Neutralizing** Immunogen Species (Isotype)

Ab Type gp120 CD4BS

Research Contact Abraham Pinter, Public Health Research Institute, Newark, NJ. pinter@phri.org

References Pinter et al. 2005 Keywords anti-idiotype

• 5145A: This study is about the V2 MAb C108g, that is typespecific and neutralizes BaL and HXB2. JR-FL is a neutralization resistant strain; modification of JRFL at V2 positions 167 and 168 (GK->DE) created a C108g epitope, and C108g could potently neutralize the modified JR-FL. The modification in V2 also increased neutralization sensitivity to V3 MABs 4117c, 2219, 2191, and 447-52D, but only had minor effects on neutralization by CD4BS MAb 5145A, and broadly neutralizing MAbs IgG1b12, 2G12, and 2F5. A modification in the NAb sensitive isolate SF162 to introduce the C108g epitope, including the introduction of a glycosylation site (160-161 KV -> NI) and 167-169 NKM -> GKV, decreased neutralization sensitivity to 5145A more than 50-fold. Pinter et al. [2005] (anti-idiotype)

**No.** 1336 MAb ID 5E HXB2 Location HIV-1 **Author Location Epitope** Neutralizing **Immunogen** Species (Isotype) human **Ab Type** gp120 CD4BS References Haynes et al. 2005

• 5E: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 5E has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

Keywords antibody binding site definition and exposure

No. 1337 **MAb ID** 8.2A **HXB2** Location HIV-1

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human

**Ab Type** gp120 C1-C4 References Haynes et al. 2005

**Keywords** antibody binding site definition and exposure • 8.2A: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. 8.2A has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

No. 1338 MAb ID E047 HXB2 Location HIV-1 **Author Location** 

**Epitope** Neutralizing

**Immunogen** 

Species (Isotype) human

Ab Type gp120 CCR5BS References Haynes et al. 2005

**Keywords** antibody binding site definition and exposure • E047: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. E047 has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

**No.** 1339 MAb ID ED10 HXB2 Location HIV-1 **Author Location Epitope Neutralizing Immunogen** Species (Isotype) human

Ab Type gp120 CCR5BS

References Haynes et al. 2005

Keywords antibody binding site definition and exposure • ED10: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. ED10 has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

**No.** 1340 MAb ID EH21 HXB2 Location HIV-1 HIV-1 Antibodies Tables HIV-1 Antibodies

Author Location
Epitope
Neutralizing
Immunogen
Species (Isotype) human

**Ab Type** gp120 C1-C4 **References** Haynes *et al.* 2005

**Keywords** antibody binding site definition and exposure
• EH21: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. EH21 has no indication of polyspecific autoreactivity. Haynes *et al.* [2005] (antibody binding site definition and exposure)

No. 1341
MAb ID F1
HXB2 Location HIV-1
Author Location
Epitope
Neutralizing
Immunogen
Species (Isotype) human

**Ab Type** gp120 CD4BS

References Haynes et al. 2005; Fujii et al. 1993

- F1: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. F1 has no indication of polyspecific autoreactivity. Haynes et al. [2005]
- F1: There is a Nef (Fujii1993) and a CD4BS (Haynes2005)
   MAb that are called F1. Fujii et al. [1993]; Haynes et al. [2005]

No. 1342 MAb ID F2A3 HXB2 Location HIV-1 Author Location

Epitope

Epitope

Neutralizing Immunogen

Species (Isotype) human

Ab Type gp120 V3

References Haynes et al. 2005

 F2A3: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. F2A3 has no indication of polyspecific autoreac-

tivity. Haynes et al. [2005] (antibody binding site definition

Keywords antibody binding site definition and exposure

and exposure)

No. 1343 MAb ID F3.9F HXB2 Location HIV-1 **Author Location** 

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype) human

**Ab Type** gp120 V3

References Haynes et al. 2005

**Keywords** antibody binding site definition and exposure • F3.9F: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. F3.9F has no indication of polyspecific autoreactivity. Haynes *et al.* [2005] (antibody binding site definition and exposure)

**No.** 1344

MAb ID LA15

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 CCR5BS

References Haynes et al. 2005

Keywords antibody binding site definition and exposure
• LA15: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. LA15 has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

**No.** 1345

MAb ID LA21

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen

Species (Isotype) human

Ab Type gp120 V3

**References** Haynes *et al.* 2005 **Keywords** antibody binding site definition and exposure

• LA21: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be dif-

polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. LA21 has no indication of polyspecific autoreactivity. Haynes *et al.* [2005] (antibody binding site definition

and exposure)

MAb ID LA28

No. 1346

HXB2 Location HIV-1

HIV-1 Antibodies HIV Antibodies Tables

Author Location Epitope

Neutralizing Immunogen

Species (Isotype) human

**Ab Type** gp120 CCR5BS **References** Haynes *et al.* 2005

Keywords antibody binding site definition and exposure

LA28: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. LA28 has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

**No.** 1347

MAb ID LF17

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

**Neutralizing** 

Immunogen

Species (Isotype) human

**Ab Type** gp120 CCR5BS **References** Haynes *et al.* 2005

**Keywords** antibody binding site definition and exposure

LF17: Of 35 Env-specific MAbs tested, only 2F5, 4E10, IgG1b12, and two CD4BS adjacent MAbs (A32 and 1.4G) and gp41 MAbs (2.2B and KU32) had binding patterns suggesting polyspecific autoreactivity, and similar reactivities may be difficult to induce with vaccines because of elimination of such autoreactivity. LF17 has no indication of polyspecific autoreactivity. Haynes et al. [2005] (antibody binding site definition and exposure)

**No.** 1348

MAb ID M2

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Species (Isotype)

Ab Type gp120 V4

References Ren et al. 2005

**Keywords** antibody binding site definition and exposure, neutralization potency

• M2: This antibody is specific for a peptide flag inserted into the V4 loop of YU-2, a neutralization resistant variant with a short V4 loop. IgG1b12 and 2F5 could neutralize both the WT YU-2 and the modified variant. The high diversity of V4 suggests it does not play a direct role in receptor binding or viral entry, yet M2, specific for the peptide insert tag, can neutralize the modified virus, demonstrating that neutralizing activity doesn't have to block funtionality of the virus. Ren et al. [2005] (antibody binding site definition and exposure, neutralization potency)

**No.** 1349

MAb ID polyclonal

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Fournier et al. 2002b

Purified B lymphocytes secret only a fraction of Ig and anti-HIV-1 Ab compared with unfractionated cells because monocytes and natural killer cells enhance both secretions by cell-to-cell contacts, involving adhesion and CD27, CD80 costimulatory molecules and IL-6 – cell-to-cell contacts and soluble factors induce maturation of activated B cells *in vitro* to allow prolonged survival and terminal differentiation. Fournier *et al.* [2002b]

**No.** 1350

MAb ID polyclonal

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human

References Fournier et al. 2002a

An early and sustained fall in plasma viral load to below detection was observed in 17 HAART responders while HIV-1 RNA remained detectable in 13 incomplete responders – HIV-1 specific Ab secretion decreased in parallel with plasma viral load – HIV-1 specific Abs became negative in only six responders, and was correlated with greater increases of CD4 T-cell counts and higher levels of HIV-specific IgA secretion at baseline – persistent immune activation may be due to residual HIV antigen. Fournier et al. [2002a]

**No.** 1351

MAb ID polyclonal

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

**Neutralizing** 

Immunogen HIV-1 infection

Species (Isotype) human

References Subbramanian et al. 2002

• Sera from 39 patients were used to study the relative prevalence of neutralizing Abs (NAbs), ADCC-Abs and enhancing Abs – 69% of the sera were positive for NAbs but only 39% could neutralize in the presence of complement – 60% had ADCC Abs – 72% mediated the enhancement of infection in the presence of complement. Subbramanian *et al.* [2002]

No. 1352

MAb ID polyclonal

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

HIV-1 Antibodies Tables HIV-1 Antibodies

Species (Isotype) human (IgA, IgG1)

References Battle-Miller et al. 2002

• In a study of HIV-1 infected women, ADCC Abs were detected in 16% (12/51) of cervicovaginal fluids, and 56% (25/45) of serum samples – 3 women had ADCC in cervical lavage fluids, but not sera, suggesting local production. Battle-Miller *et al.* [2002]

**No.** 1353

MAb ID polyclonal

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgA1, IgA2, IgM)

References Wu & Jackson 2002

• IgA1 accounted for the majority of anti-HIV-1 IgA in the saliva in HIV-1 infected individuals – there was no anti-gp41 IgA in saliva, in contrast to plasma – lower levels of IgA and IgM were found in saliva than in plasma. Wu & Jackson [2002]

**No.** 1354

MAb ID polyclonal

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype B

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Hioe et al. 1997a

• Four primary isolates showed distinct patterns of sensitivity to neutralization by polyclonal sera or plasma and MAbs – BZ167 was the only isolate inhibited by all polyclonal sera and plasma tested, and was also neutralized by 8/17 MAbs, in particular anti-V3 loop (419-D, 447-52D, 782-D, and 838-D), anti-CD4bd (559/64-D, 654-D and 830-D and a cluster II of gp41 directed MAb (98-6) – isolates 92HT593 and 91US056 were neutralized by V3 loop (419-D, and 447-52D)and cluster II gp41 (98-6) MAbs at higher concentrations – US4 was neutralized by some of the polyclonal sera/plasma tested and not at all by MAbs individually or by a cocktail of ten MAbs consisting of 419-D, 447-52D, 782-D, 838-D, 559/64-D, 654-D, 450-D, 670-D, 1281-D and 98-6. Hioe *et al.* [1997a]

**No.** 1355

MAb ID polyclonal

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Neutralizing no

Immunogen HIV-1 infection

Species (Isotype) human (IgA, IgG)

References Oelemann et al. 2002

 A urine based commercial EIA kit from Calypte Biomedical Corporation, Berkeley, CA was found to work well as a primary screening for HIV in Brazilian samples – 76 HIV+ samples were correctly identified (100% sensitivity), and 278/284 negative samples 97.9% specificity. Oelemann *et al.* [2002] **No.** 1356

MAb ID polyclonal

**HXB2** Location HIV-1

Author Location HIV-1

**Epitope** 

Neutralizing no

**Immunogen** HIV-1 infection

Species (Isotype) human (IgE)

References Pellegrino et al. 2002; Secord et al. 1996

- Pediatric long term survivors (LTS) have been found to carry HIV-1 specific IgE serum from these children inhibit HIV-1 production in culture, but this inhibition did not seem to be due to neutralization, rather due to a cytoxic event serum lost the HIV-1 inhibitory effect when depleted of IgE. Pellegrino *et al.* [2002]
- HIV-specific IgE found in clinically healthy HIV-1 infected children. Second et al. [1996]

**No.** 1357

MAb ID polyclonal

**HXB2** Location HIV-1

Author Location gp120 and p55

**Epitope** 

Neutralizing no

Immunogen vaccine

Vector/Type: vaccinia Strain: B clade 89.6 HIV component: Env, Gag-Pol Adjuvant: E. coli mutant heat labile enterotoxin (LT-R72)

Species (Isotype) macaque

References Ambrose et al. 2003

Keywords genital and mucosal immunity

Systemic priming with rVVs expressing HIV-1 Env and SHIV Gag-Pol followed by intragastric and intranasal mucosal boosting of LT(R192G) and aldrithiol-2 (AT-2)-inactiviated SHIV induced SHIV-specific IgA and IgG plasma and mucosal Abs. Viral loads in vaccinated animals were reduced after vaginal challenge with SHIV 89.6. Ambrose *et al.* [2003] (genital and mucosal immunity)

No. 1358

MAb ID polyclonal

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype B

Neutralizing P

Immunogen HIV-1 infection

Species (Isotype) human

References Binley et al. 2004

**Keywords** subtype comparisons, variant cross-recognition or cross-neutralization

• 93 viruses from different clades were tested for their neutralization cross-reactivity using a panel of HIV monoclonal antibodies, and a plasma from an HIV-1 + donor infected with a B clade virus. The plasma antibodies broadly neutralized viruses from many clades, with a slight preference for B clade. Binley *et al.* [2004] (variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 1359

HIV-1 Antibodies HIV Antibodies Tables

MAb ID polyclonal

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype multiple

Neutralizing

Immunogen vaccine

Vector/Type: peptide Strain: natural vari-

ants HIV component: gp140

Species (Isotype) rabbit

**Ab Type** RT thumb domain **References** Dong *et al.* 2005

**Keywords** vaccine antigen design, variant crossrecognition or cross-neutralization

2F5 recognizes the epitope ELDKWA, but does not neutralize viruses carrying the commonly found mutated epitope variants: ELDEWA, ELDSWA, ELDNWA, ELDQWA, ELDTWA, or ELNKWA. Peptide cocktails containing ELDKWA, ELNKWA, ELDEWA, and ELEKWA elicit polyclonal antibodies in rabbits that can bind to all of the natural variants that are escape variants for 2F5 expressed in gp41 via WB, as well as ELDERWA. Dong et al. [2005] (vaccine antigen design, variant cross-recognition or cross-neutralization)

**No.** 1360

MAb ID polyclonal

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype multiple

**Neutralizing** 

Immunogen vaccine

Vector/Type: peptide HIV component: gp41 Adjuvant: Complete Freund's Adjuvant (CFA), Incomplete Freund's Adjuvant (IFA)

Species (Isotype) rabbit

References Dong et al. 2005

**Keywords** escape, vaccine antigen design, variant crossrecognition or cross-neutralization

• 2F5 recognizes the epitope ELDKWA, but does not neutralize viruses carrying the commonly found mutated epitope, ELDEWA, ELDSWA, ELDNWA, ELDQWA, ELDTWA, or ELNKWA. Peptide cocktails containing ELDKWA, ELNKWA, ELDEWA, and ELEKWA, elicit polyclonal antibodies in rabbits that can bind to all of the natural variants that are escape variants for 2F5 expressed in gp41 via WB, as well as ELDERWA. Dong *et al.* [2005] (vaccine antigen design, variant cross-recognition or cross-neutralization, escape)

**No.** 1361

MAb ID polyclonal

HXB2 Location HIV-1

**Author Location** 

**Epitope** 

Subtype A, B

Neutralizing L

Immunogen HIV-1 infection

Species (Isotype) human

Ab Type gp120 V3

References Krachmarov et al. 2005

Country Cameroon

**Keywords** antibody binding site definition and exposure, subtype comparisons, variant cross-recognition or cross-neutralization

• Sera from 23 subtype A infected individuals from Cameroon have antibodies that react strongly with subtype A and subtype B V3 loops in fusion proteins, and neutralize SF162 pseudotypes, while sera from 47 subtype B infected individuals reacted only with subtype B. The sera from Cameroon do not neutralize primary A or B isolates, due to indirect masking by the V1/V2 domain rather than due to loss of the target epitope. Krachmarov *et al.* [2005] (antibody binding site definition and exposure, variant cross-recognition or cross-neutralization, subtype comparisons)

**No.** 1362

MAb ID polyclonal

HXB2 Location HIV-1

Author Location (HXB2)

**Epitope** 

Subtype B

Neutralizing yes

Immunogen vaccine

Vector/Type: vesicular stomatitis virus (VSV) with protein boost Strain: B clade HXB2 HIV component: gp41 MPER Adjuvant: Complete Freund's Adjuvant (CFA), Incomplete Freund's Adjuvant (IFA)

Species (Isotype) rabbit

References Luo et al. 2006

Keywords vaccine antigen design

• gp41 and p15E of the porcine endogenous retrovirus (PERV) share structural and functional similarities, and epitopes in the membrane proximal region of p15E are able to elicit NAbs upon immunization with soluble p15E. Rabbits immunized with a VSV recombinant expressing an HIV-1 membrane-proximal external region (MPER) fused to PERV p15E, with a fusion p15E-HIV MPER protein boost, elicted HIV specific NAbs in 3/9 rabbits each for two different constructs, with and without the E2 region of p15E. Luo et al. [2006] (vaccine antigen design)

**No.** 1363

MAb ID polyclonal

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype multiple

Neutralizing

Immunogen HIV-1 infection

Species (Isotype) human (IgG)

References Parekh & McDougal 2005

**Keywords** acute/early infection, assay development, assay standardization/improvement

• This paper describes IgG-Capture BED-EIA, which detects the increasing proportion of HIV-1-IgG relative to total IgG and can be used to detect early infection and incidence data in cross-sectional and sentinel surveillance studies, and is robust for use with multiple HIV-1 subtypes. Parekh & McDougal [2005]

HIV Antibodies Tables HIV-1 Antibodies

(assay development, acute/early infection, assay standardization/improvement)

**No.** 1364

MAb ID polyclonal

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: fixed fusion-intermediate Strain: B clade US005.11, FASH isolate

HIV component: Env

Species (Isotype) mouse

References Zipeto et al. 2005

Keywords co-receptor, vaccine antigen design

• HIV-1 fusion complexes were prepared from cell lines expressing R5 HIV-1 gp120/gp41 and CD4-CCR5. Fusion complexes were prepared at different temperatures (21, 30 or 37 degrees C) with different fixative combinations, and used to immunize mice. Complexes prepared at 37 degrees were the most immunogenic, suggesting that fixation of multiple conformation intermediates may by helpful. Neutralizing Abs were raised against both R5 (strain BaL) and X4 (strain 213) viruses. Zipeto et al. [2005] (co-receptor, vaccine antigen design)

**No.** 1365

MAb ID polyclonal

**HXB2** Location HIV-1

**Author Location** 

**Epitope** 

Subtype B

Neutralizing no

Immunogen vaccine

Vector/Type: protein Strain: B clade ADA HIV component: Env fragments in a pre-fusion state trimer Adjuvant: Complete Freund's Adjuvant (CFA), Incomplete Freund's Adjuvant (IFA)

Species (Isotype) rabbit

References Qiao et al. 2005

**Keywords** antibody binding site definition and exposure, vaccine antigen design, vaccine-specific epitope characteristics

 A gp140 prefusion state trimer composed of gp41 truncated at Lys665, and gp120 C1 and C5 (topless gp140), was engineered and used to immune rabbits. No NAbs were raised, although the polyclonal sera recognized many regions of the truncated Env. Qiao et al. [2005] (antibody binding site definition and exposure, vaccine antigen design, vaccine-specific epitope characteristics)

**No.** 1366

MAb ID polyclonal

**HXB2 Location** HIV-1

**Author Location** 

**Epitope** 

Subtype B

Neutralizing

Immunogen vaccine

Vector/Type: DNA prime with modified vaccinia Ankara (MVA) boost HIV component:

Env, Gag-Pol

Species (Isotype) macaque

References Sadagopal et al. 2005

Keywords vaccine-induced epitopes

• 22/23 macaques vaccinated with a DNA Gag-Pol\_Env prime and vaccinia virus Ankara boost controlled SHIV viremia until euthanasia at 200 weeks post-challenge. All animals had low or undetectable viral loads, normal CD4 counts, and high titers of neutralizing antibodies. Most animals recognized 2 CD8 epitopes and 1 CD4 epitope, with up to 3 CD8 and 5 CD4 epitopes. Most T-cell epitopes were in Gag, though some were in Env. Sadagopal et al. [2005] (vaccine-induced epitopes)

# **IV-D**

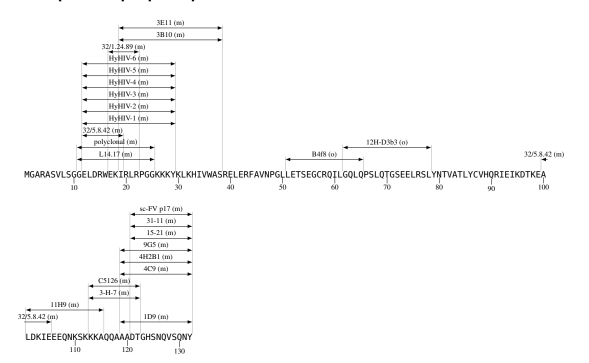
# Maps of MAb Locations Plotted by Protein

Linear epitopes less than of 21 amino acids or less are shown with their antibody ID and the experimental species.

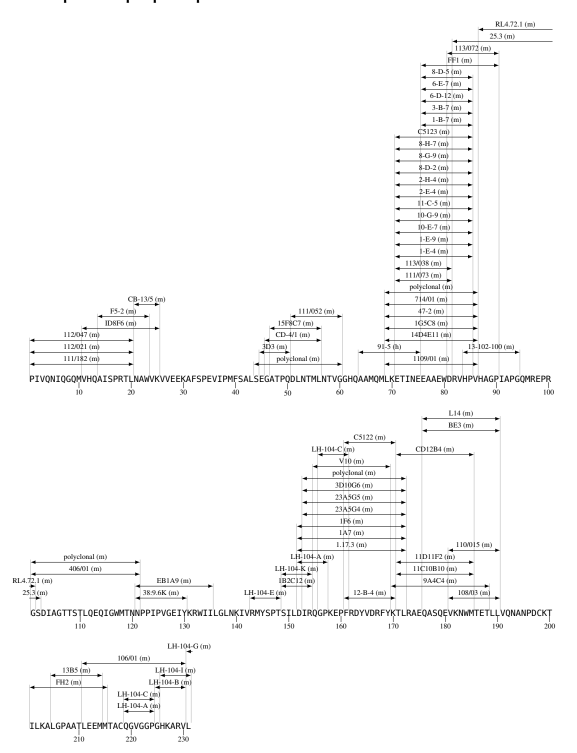
Key	Species
h	human
p	non-human primate
m	murine
O	other

**Table IV-D.1:** The species that the epitope was generated in and derived from.

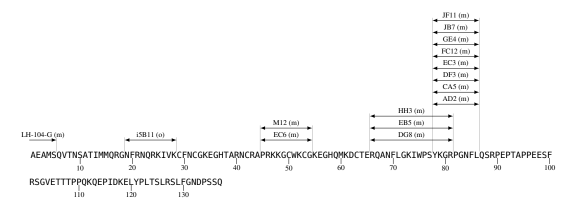
# IV-D-1 p17 Ab Epitope Map



# IV-D-2 p24 Ab Epitope Map



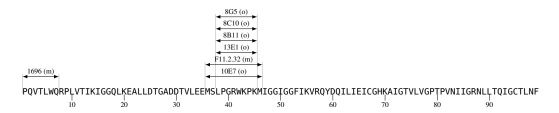
#### IV-D-3 p2p7p1p6 Ab Epitope Map



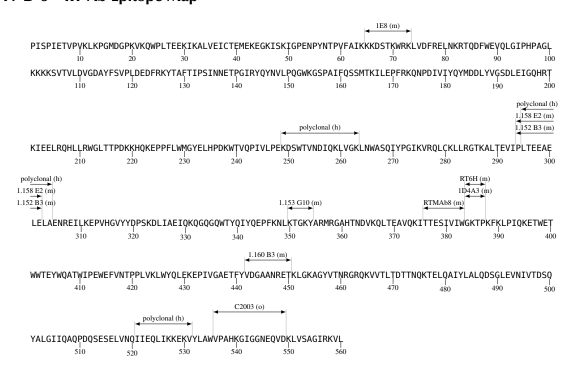
#### IV-D-4 Gag/Pol TF Ab Epitope Map

FFREDLAFLOGKAREFSSEQTRANSPTRRELQVWGRDNNSPSEAGADROGTVSFNF

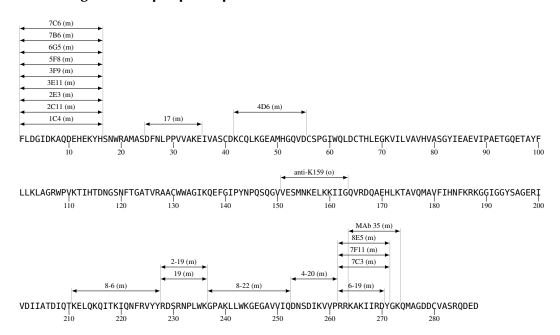
#### IV-D-5 Protease Ab Epitope Map



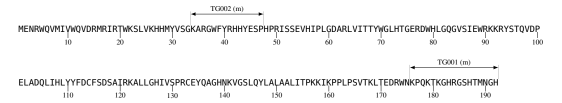
#### IV-D-6 RT Ab Epitope Map



#### IV-D-7 Integrase Ab Epitope Map



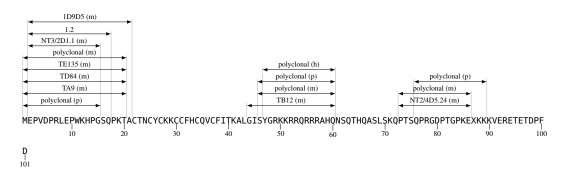
#### IV-D-8 Vif Ab Epitope Map



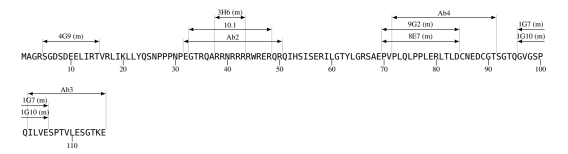
#### IV-D-9 Vpr Ab Epitope Map

MEQAPEDQGPQREPHNEWTLELLEELKNEAVRHFPRIWLHGLGQHIYETYGDTWAGVEAIIRILQQLLFIHFRIGCRHSRIGVTRQRRARNGASRS

#### IV-D-10 Tat Ab Epitope Map

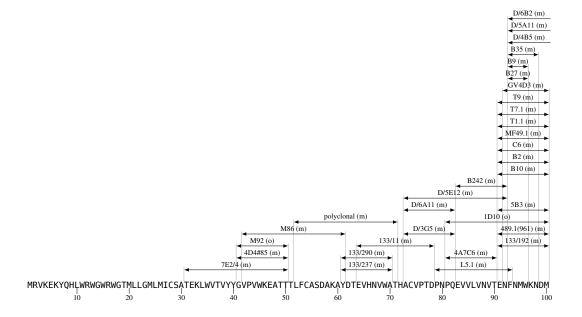


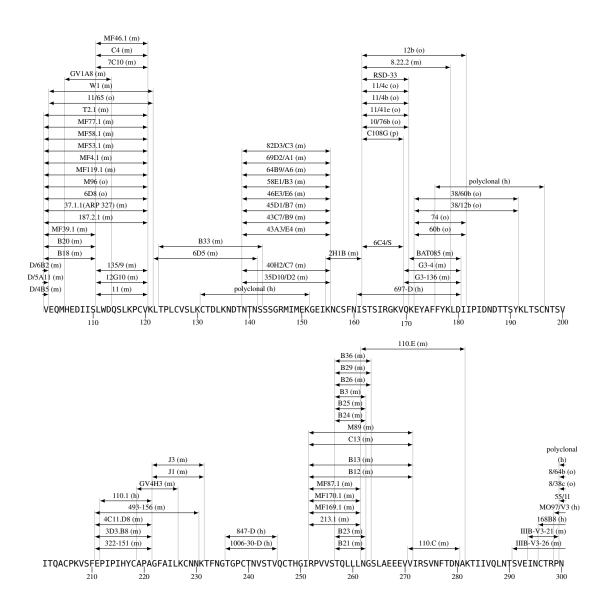
# IV-D-11 Rev Ab Epitope Map

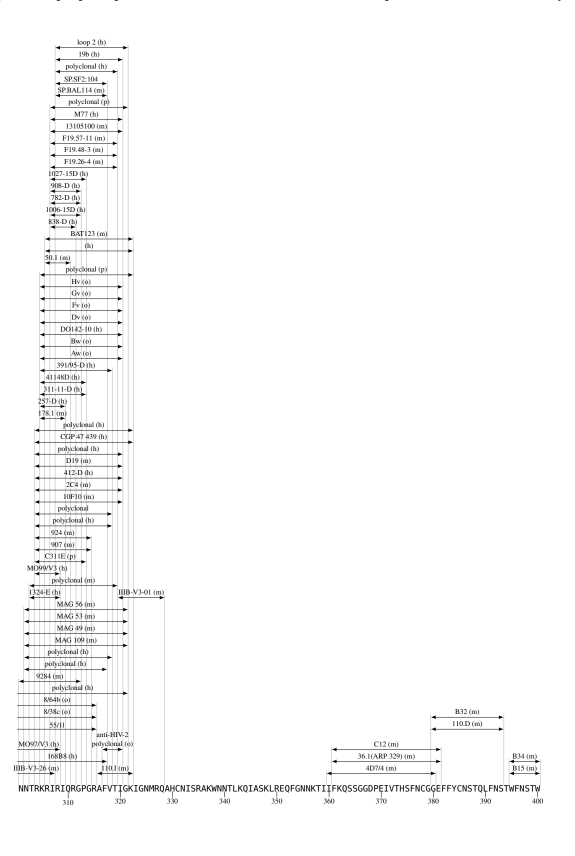


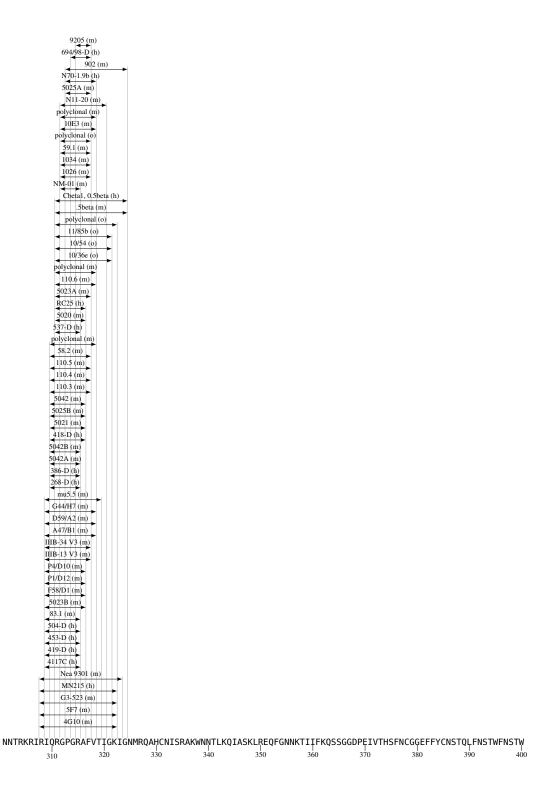
#### IV-D-12 Vpu Ab Epitope Map

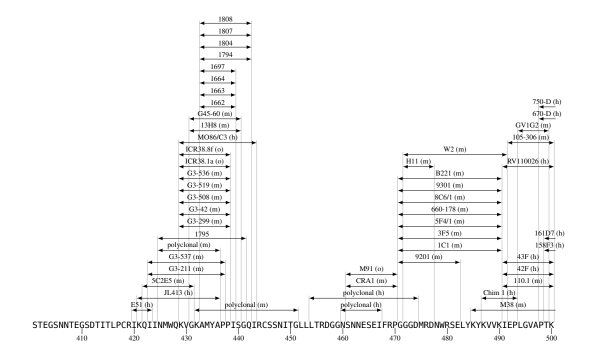
# IV-D-13 gp160 Ab Epitope Map

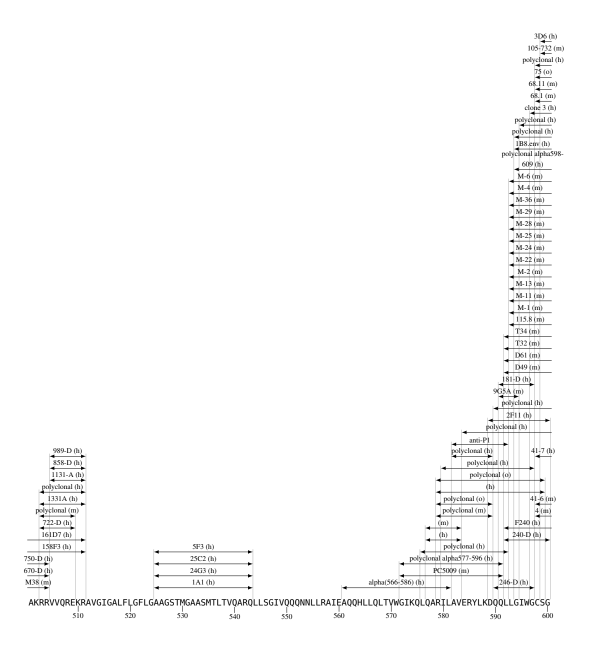


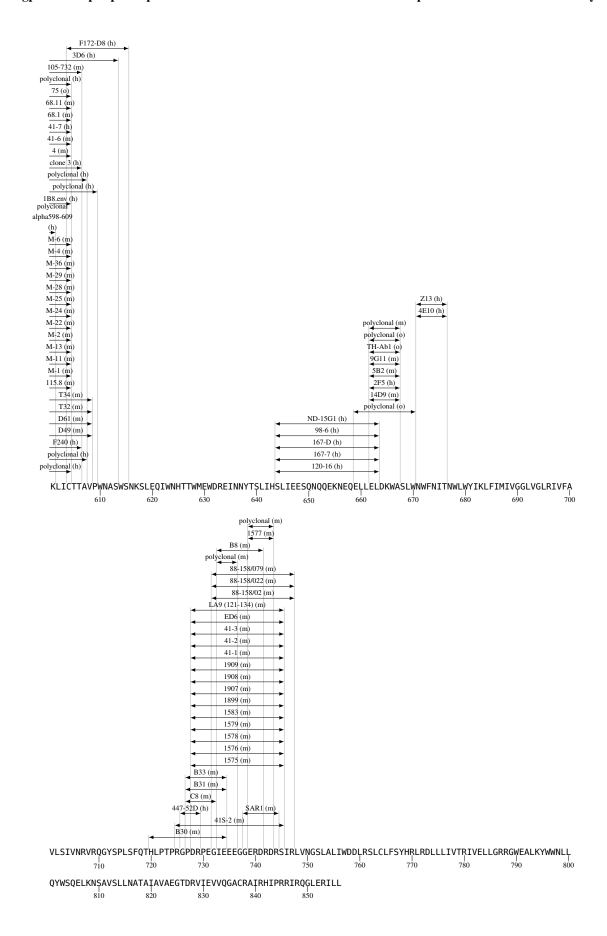




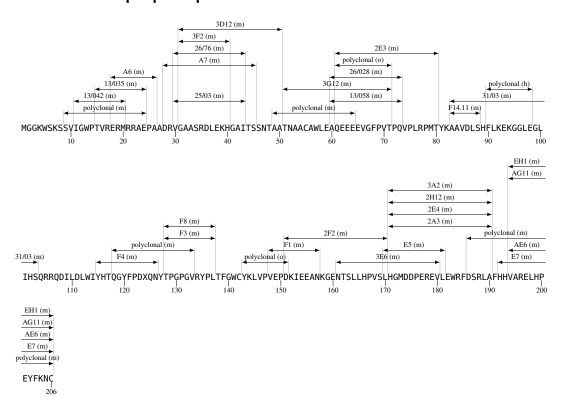








# IV-D-14 Nef Ab Epitope Map



# Part V HIV Immunology References

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